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(57) Abstract

The sequences of 5' ESTs derived from mRNAs encoding secreted proteins are disclosed. The 5' ESTs may be to obtain cDNAs and genomic DNAs corresponding to the 5' ESTs. The 5' ESTs may also be used in diagnostic, forensic, gene therapy, and chromosome mapping procedures. Upstream regulatory sequences may also be obtained using the 5' ESTs. The 5' ESTs may also be used to design expression vectors and secretion vectors.

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5' ESTs FOR SECRETED PROTEINS EXPRESSED IN PROSTATE

Background of the Invention

The estimated 50,000-100,000 genes scattered along the human chromosomes offer tremendous promise for the understanding, diagnosis, and treatment of human diseases. In addition, probes capable of specifically hybridizing to loci distributed throughout the human genome find applications in the construction of high resolution chromosome maps and in the identification of individuals.

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In the past, the characterization of even a single human gene was a painstaking process, requiring years of effort. Recent developments in the areas of cloning vectors, DNA sequencing, and computer technology have merged to greatly accelerate the rate at which human genes can be isolated, sequenced, mapped, and characterized. Cloning vectors such as yeast artificial chromosomes (YACs) and bacterial artificial chromosomes (BACs) are able to accept DNA inserts ranging from 300 to 1000 kilobases (kb) or 100-400 kb in length respectively, thereby facilitating the manipulation and ordering of DNA sequences distributed over great distances on the human chromosomes. Automated DNA sequencing machines permit the rapid sequencing of human genes. Bioinformatics software enables the comparison of nucleic acid and protein sequences, thereby assisting in the characterization of human gene products.

Currently, two different approaches are being pursued for identifying and characterizing the genes distributed along the human genome. In one approach, large fragments of genomic DNA are isolated, cloned, and sequenced. Potential open reading frames in these genomic sequences are identified using bioinformatics software. However, this approach entails sequencing large stretches of human DNA which do not encode proteins in order to find the protein encoding sequences scattered throughout the genome. In addition to requiring extensive sequencing, the bioinformatics software may mischaracterize the genomic sequences obtained. Thus, the software may produce false positives in which noncoding DNA is mischaracterized as coding DNA or false negatives in which coding DNA is mischaracterized as non-coding DNA.

An alternative approach takes a more direct route to identifying and characterizing human genes. In this approach, complementary DNAs (cDNAs) are synthesized from isolated messenger RNAs (mRNAs) which encode human proteins. Using this approach,

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sequencing is only performed on DNA which is derived from protein coding portions of the genome. Often, only short stretches of the cDNAs are sequenced to obtain sequences called expressed sequence tags (ESTs). The ESTs may then be used to isolate or purify extended cDNAs which include sequences adjacent to the EST sequences. The extended cDNAs may contain all of the sequence of the EST which was used to obtain them or only a portion of the sequence of the EST which was used to obtain them. In addition, the extended cDNAs may contain the full coding sequence of the gene from which the EST was derived or, alternatively, the extended cDNAs may include portions of the coding sequence of the gene from which the EST was derived. It will be appreciated that there may be several extended cDNAs which include the EST sequence as a result of alternate splicing or the activity of alternative promoters.

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In the past, these short EST sequences were often obtained from oligo-dT primed cDNA libraries. Accordingly, they mainly corresponded to the 3' untranslated region of the mRNA. In part, the prevalence of EST sequences derived from the 3' end of the mRNA is a result of the fact that typical techniques for obtaining cDNAs are not well suited for isolating cDNA sequences derived from the 5' ends of mRNAs. (Adams *et al.*, *Nature* 377:3-174, 1996; Hillier *et al.*, *Genome Res.* 6:807-828, 1996).

In addition, in those reported instances where longer cDNA sequences have been obtained, the reported sequences typically correspond to coding sequences and do not include the full 5' untranslated region of the mRNA from which the cDNA is derived. Such incomplete sequences may not include the first exon of the mRNA, particularly in situations where the first exon is short. Furthermore, they may not include some exons, often short ones, which are located upstream of splicing sites. Thus, there is a need to obtain sequences derived from the 5' ends of mRNAs.

While many sequences derived from human chromosomes have practical applications, approaches based on the identification and characterization of those chromosomal sequences which encode a protein product are particularly relevant to diagnostic and therapeutic uses. Of the 50,000-100,000 protein coding genes, those genes encoding proteins which are secreted from the cell in which they are synthesized, as well as the secreted proteins themselves, are particularly valuable as potential therapeutic agents. Such proteins are often

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involved in cell to cell communication and may be responsible for producing a clinically relevant response in their target cells.

In fact, several secretory proteins, including tissue plasminogen activator, G-CSF, GM-CSF, erythropoietin, human growth hormone, insulin, interferon-α, interferon-β, interferon-γ, and interleukin-2, are currently in clinical use. These proteins are used to treat a wide range of conditions, including acute myocardial infarction, acute ischemic stroke, anemia, diabetes, growth hormone deficiency, hepatitis, kidney carcinoma, chemotherapy induced neutropenia and multiple sclerosis. For these reasons, extended cDNAs encoding secreted proteins or portions thereof represent a particularly valuable source of therapeutic agents. Thus, there is a need for the identification and characterization of secreted proteins and the nucleic acids encoding them.

In addition to being therapeutically useful themselves, secretory proteins include short peptides, called signal peptides, at their amino termini which direct their secretion. These signal peptides are encoded by the signal sequences located at the 5' ends of the coding sequences of genes encoding secreted proteins. Because these signal peptides will direct the extracellular secretion of any protein to which they are operably linked, the signal sequences may be exploited to direct the efficient secretion of any protein by operably linking the signal sequences to a gene encoding the protein for which secretion is desired. In addition, portions of signal sequences may also be used to direct the intracellular import of a peptide or protein of interest. This may prove beneficial in gene therapy strategies in which it is desired to deliver a particular gene product to cells other than the cell in which it is produced. Signal sequences encoding signal peptides also find application in simplifying protein purification techniques. In such applications, the extracellular secretion of the desired protein greatly facilitates purification by reducing the number of undesired proteins from which the desired protein must be selected. Thus, there exists a need to identify and characterize the 5' portions of the genes for secretory proteins which encode signal peptides.

Public information on the number of human genes for which the promoters and upstream regulatory regions have been identified and characterized is quite limited. In part, this may be due to the difficulty of isolating such regulatory sequences. Upstream regulatory sequences such as transcription factor binding sites are typically too short to be utilized as probes for isolating promoters from human genomic libraries. Recently, some approaches

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have been developed to isolate human promoters. One of them consists of making a CpG island library (Cross, et al., Nature Genetics 6: 236-244, 1994). The second consists of isolating human genomic DNA sequences containing SpeI binding sites by the use of SpeI binding protein. (Mortlock et al., Genome Res. 6:327-335, 1996). Both of these approaches have their limits due to a lack of specificity or of comprehensiveness.

The present 5' ESTs may be used to efficiently identify and isolate upstream regulatory regions which control the location, developmental stage, rate, and quantity of protein synthesis, as well as the stability of the mRNA. (Theil, *BioFactors* 4:87-93, 1993). Once identified and characterized, these regulatory regions may be utilized in gene therapy or protein purification schemes to obtain the desired amount and locations of protein synthesis or to inhibit, reduce, or prevent the synthesis of undesirable gene products.

In addition, ESTs containing the 5' ends of secretory protein genes may include sequences useful as probes for chromosome mapping and the identification of individuals. Thus, there is a need to identify and characterize the sequences upstream of the 5' coding sequences of genes encoding secretory proteins.

Summary of the Invention

The present invention relates to purified, isolated, or recombinant ESTs which include sequences derived from the authentic 5' ends of their corresponding mRNAs. The term "corresponding mRNA" refers to the mRNA which was the template for the cDNA synthesis which produced the 5' EST. These sequences will be referred to hereinafter as "5' ESTs." As used herein, the term "purified" does not require absolute purity; rather, it is intended as a relative definition. Individual 5' EST clones isolated from a cDNA library have been conventionally purified to electrophoretic homogeneity. The sequences obtained from these clones could not be obtained directly either from the library or from total human DNA. The cDNA clones are not naturally occurring as such, but rather are obtained via manipulation of a partially purified naturally occurring substance (messenger RNA). The conversion of mRNA into a cDNA library involves the creation of a synthetic substance (cDNA) and pure individual cDNA clones can be isolated from the synthetic library by clonal selection. Thus, creating a cDNA library from messenger RNA and subsequently isolating individual clones from that library results in an approximately 10⁴-10⁶ fold purification of the native message.

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Purification of starting material or natural material to at least one order of magnitude, preferably two or three orders, and more preferably four or five orders of magnitude is expressly contemplated.

As used herein, the term "isolated" requires that the material be removed from its original environment (e.g., the natural environment if it is naturally occurring). For example, a naturally-occurring polynucleotide present in a living animal is not isolated, but the same polynucleotide, separated from some or all of the coexisting materials in the natural system, is isolated.

As used herein, the term "recombinant" means that the 5' EST is adjacent to "backbone" nucleic acid to which it is not adjacent in its natural environment. Additionally, to be "enriched" the 5' ESTs will represent 5% or more of the number of nucleic acid inserts in a population of nucleic acid backbone molecules. Backbone molecules according to the present invention include nucleic acids such as expression vectors, self-replicating nucleic acids, viruses, integrating nucleic acids, and other vectors or nucleic acids used to maintain or manipulate a nucleic acid insert of interest. Preferably, the enriched 5' ESTs represent 15% or more of the number of nucleic acid inserts in the population of recombinant backbone molecules. More preferably, the enriched 5' ESTs represent 50% or more of the number of nucleic acid inserts in the population of recombinant backbone molecules. In a highly preferred embodiment, the enriched 5' ESTs represent 90% or more of the number of nucleic acid inserts in the population of recombinant backbone molecules.

"Stringent", moderate," and "low" hybridization conditions are as defined in Example 29.

Unless otherwise indicated, a "complementary" sequence is fully complementary.

Thus, 5' ESTs in cDNA libraries in which one or more 5' ESTs make up 5% or more of the number of nucleic acid inserts in the backbone molecules are "enriched recombinant 5' ESTs" as defined herein. Likewise, 5' ESTs in a population of plasmids in which one or more 5' EST of the present invention have been inserted such that they represent 5% or more of the number of inserts in the plasmid backbone are " enriched recombinant 5' ESTs" as defined herein. However, 5' ESTs in cDNA libraries in which 5' ESTs constitute less than 5% of the number of nucleic acid inserts in the population of backbone molecules, such as libraries in

6

which backbone molecules having a 5' EST insert are extremely rare, are not "enriched recombinant 5' ESTs."

In particular, the present invention relates to 5' ESTs which are derived from genes encoding secreted proteins. As used herein, a "secreted" protein is one which, when expressed in a suitable host cell, is transported across or through a membrane, including transport as a result of signal peptides in its amino acid sequence. "Secreted" proteins include without limitation proteins secreted wholly (e.g. soluble proteins), or partially (e.g. receptors) from the cell in which they are expressed. "Secreted" proteins also include without limitation proteins which are transported across the membrane of the endoplasmic reticulum.

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Such 5' ESTs include nucleic acid sequences, called signal sequences, which encode signal peptides which direct the extracellular secretion of the proteins encoded by the genes from which the 5' ESTs are derived. Generally, the signal peptides are located at the amino termini of secreted proteins.

Secreted proteins are translated by ribosomes associated with the "rough" endoplasmic reticulum. Generally, secreted proteins are co-translationally transferred to the membrane of the endoplasmic reticulum. Association of the ribosome with the endoplasmic reticulum during translation of secreted proteins is mediated by the signal peptide. The signal peptide is typically cleaved following its co-translational entry into the endoplasmic reticulum. After delivery to the endoplasmic reticulum, secreted proteins may proceed through the Golgi apparatus. In the Golgi apparatus, the proteins may undergo post-translational modification before entering secretory vesicles which transport them across the cell membrane.

The 5' ESTs of the present invention have several important applications. For example, they may be used to obtain and express cDNA clones which include the full protein coding sequences of the corresponding gene products, including the authentic translation start sites derived from the 5' ends of the coding sequences of the mRNAs from which the 5' ESTs are derived. These cDNAs will be referred to hereinafter as "full length cDNAs." These cDNAs may also include DNA derived from mRNA sequences upstream of the translation start site. The full length cDNA sequences may be used to express the proteins corresponding to the 5' ESTs. As discussed above, secreted proteins are therapeutically important. Thus, the proteins expressed from the cDNAs may be useful in treating or

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controlling a variety of human conditions. The 5' ESTs may also be used to obtain the corresponding genomic DNA. The term "corresponding genomic DNA" refers to the genomic DNA which encodes the mRNA from which the 5' EST was derived.

Alternatively, the 5' ESTs may be used to obtain and express extended cDNAs encoding portions of the secreted protein. The portions may comprise the signal peptides of the secreted proteins or the mature proteins generated when the signal peptide is cleaved off. The portions may also comprise polypeptides having at least 10 consecutive amino acids encoded by the extended cDNAs or full length cDNAs. Alternatively, the portions may comprise at least 15 consecutive amino acids encoded by the extended cDNAs or full length cDNAs. In some embodiments, the portions may comprise at least 25 consecutive amino acids encoded by the extended cDNAs or full length cDNAs. In other embodiments, the portions may comprise at least 40 amino acids encoded by the extended cDNAs or full length cDNAs.

Antibodies which specifically recognize the entire secreted proteins encoded by the extended cDNAs, full length cDNAs, or fragments thereof having at least 10 consecutive amino acids, at least 15 consecutive amino acids, at least 25 consecutive amino acids, or at least 40 consecutive amino acids may also be obtained as described below. Antibodies which specifically recognize the mature protein generated when the signal peptide is cleaved may also be obtained as described below. Similarly, antibodies which specifically recognize the signal peptides encoded by the extended cDNAs or full length cDNAs may also be obtained.

In some embodiments, the extended cDNAs obtained using the 5' ESTs include the signal sequence. In other embodiments, the extended cDNAs obtained using the 5' ESTs may include the full coding sequence for the mature protein (i.e. the protein generated when the signal polypeptide is cleaved off). In addition, the extended cDNAs obtained using the 5' ESTs may include regulatory regions upstream of the translation start site or downstream of the stop codon which control the amount, location, or developmental stage of gene expression.

As discussed above, secreted proteins are therapeutically important. Thus, the proteins expressed from the extended cDNAs or full length cDNAs obtained using the 5' ESTs may be useful in treating or controlling a variety of human conditions.

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The 5' ESTs (or cDNAs or genomic DNAs obtained therefrom) may be used in forensic procedures to identify individuals or in diagnostic procedures to identify individuals having genetic diseases resulting from abnormal expression of the genes corresponding to the 5' ESTs. In addition, the present invention is useful for constructing a high resolution map of the human chromosomes.

The present invention also relates to secretion vectors capable of directing the secretion of a protein of interest. Such vectors may be used in gene therapy strategies in which it is desired to produce a gene product in one cell which is to be delivered to another location in the body. Secretion vectors may also facilitate the purification of desired proteins.

The present invention also relates to expression vectors capable of directing the expression of an inserted gene in a desired spatial or temporal manner or at a desired level. Such vectors may include sequences upstream of the 5' ESTs, such as promoters or upstream regulatory sequences.

Finally, the present invention may also be used for gene therapy to control or treat genetic diseases. Signal peptides may also be fused to heterologous proteins to direct their extracellular secretion.

Bacterial clones containing Bluescript plasmids having inserts containing the 5' ESTs of the present invention (SEQ ID NOs: 38-315 are presently stored at 80°C in 4% (v/v) glycerol in the inventor's laboratories under the designations listed next to the SEQ ID NOs in II). The inserts may be recovered from the deposited materials by growing the appropriate clones on a suitable medium. The Bluescript DNA can then be isolated using plasmid isolation procedures familiar to those skilled in the art such as alkaline lysis minipreps or large scale alkaline lysis plasmid isolation procedures. If desired the plasmid DNA may be further enriched by centrifugation on a cesium chloride gradient, size exclusion chromatography, or anion exchange chromatography. The plasmid DNA obtained using these procedures may then be manipulated using standard cloning techniques familiar to those skilled in the art. Alternatively, a PCR can be done with primers designed at both ends of the EST insertion. The PCR product which corresponds to the 5' EST can then be manipulated using standard cloning techniques familiar to those skilled in the art.

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One aspect of the present invention is a purified or isolated nucleic acid having the sequence of one of SEQ ID NOs: 38-315 or having a sequence complementary thereto. In one embodiment, the nucleic acid is recombinant.

Another aspect of the present invention is a purified or isolated nucleic acid comprising at least 10 consecutive bases of the sequence of one of SEQ ID NOs: 38-315 or one of the sequences complementary thereto.

Yet another aspect of the present invention is a purified or isolated nucleic acid comprising at least 15 consecutive bases of one of the sequences of SEQ ID NOs: 38-315 or one of the sequences complementary thereto. In one embodiment, the nucleic acid is recombinant.

A further aspect of the present invention is a purified or isolated nucleic acid of at least 15 bases capable of hybridizing under stringent conditions to the sequence of one of SEQ ID NOs: 38-315 or one of the sequences complementary to the sequences of SEQ ID NOs: 38-315. In one embodiment, the nucleic acid is recombinant.

Another aspect of the present invention is a purified or isolated nucleic acid encoding a human gene product, said human gene product having a sequence partially encoded by one of the sequences of SEQ ID NO: 38-315.

Still another aspect of the present invention is a method of making a cDNA encoding a human secretory protein, said human secretory protein being partially encoded by one of SEQ ID NOs 38-315, comprising the steps of contacting a collection of mRNA molecules from human cells with a primer comprising at least 15 consecutive nucleotides of a sequence complementary to one of SEQ ID NOs: 38-315; hybridizing said primer to an mRNA in said collection that encodes said protein; reverse transcribing said hybridized primer to make a first cDNA strand from said mRNA; making a second cDNA strand complementary to said first cDNA strand; and isolating the resulting cDNA encoding said protein comprising said first cDNA strand and said second cDNA strand.

Another aspect of the invention is an isolated or purified cDNA encoding a human secretory protein, said human secretory protein comprising the protein encoded by one of SEQ ID NOs 38-315 or a fragment thereof of at least 10 amino acids, said cDNA being obtainable by the method described in the preceding paragraph. In one embodiment, the

cDNA comprises the full protein coding sequence of said protein which sequence is partially included in one of the sequences of SEQ ID NOs: 38-315.

Another aspect of the present invention is a method of making a cDNA encoding a human secretory protein that is partially encoded by one of SEQ ID NOs 38-315, comprising the steps of obtaining a cDNA comprising one of the sequences of SEQ ID NOs: 38-315; contacting said cDNA with a detectable probe comprising at least 15 consecutive nucleotides of said sequence of SEQ ID NO: 38-315 or a sequence complementary thereto under conditions which permit said probe to hybridize to said cDNA; identifying a cDNA which hybridizes to said detectable probe; and isolating said cDNA which hybridizes to said probe.

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Another aspect of the present invention is an isolated or purified cDNA encoding a human secretory protein, said human secretory protein comprising the protein encoded by one of SEQ ID NOs 38-315 or a fragment thereof of at least 10 amino acids, said cDNA being obtainable by the method described in the preceding paragraph. In one embodiment, the cDNA comprises the full protein coding sequence partially included in one of the sequences of SEQ ID NOs: 38-315.

Another aspect of the present invention is a method of making a cDNA comprising one of the sequence of SEQ ID NOs: 38-315, comprising the steps of contacting a collection of mRNA molecules from human cells with a first primer capable of hybridizing to the polyA tail of said mRNA; hybridizing said first primer to said polyA tail; reverse transcribing said mRNA to make a first cDNA strand; making a second cDNA strand complementary to said first cDNA strand using at least one primer comprising at least 15 nucleotides of one of the sequences of SEQ ID NOs 38-315; and isolating the resulting cDNA comprising said first cDNA strand and said second cDNA strand.

Another aspect of the present invention is an isolated or purified cDNA encoding a human secretory protein, said human secretory protein comprising the protein encoded by one of SEQ ID NOs 38-315 or a fragment thereof of at least 10 amino acids, said cDNA being obtainable by the method described in the preceding paragraph. In one embodiment, the cDNA comprises the full protein coding sequence partially included in one of the sequences of SEQ ID NOs: 38-315.

In one embodiment of the method described in the two paragraphs above, the second cDNA strand is made by contacting said first cDNA strand with a first pair of primers, said

first pair of primers comprising a second primer comprising at least 15 consecutive nucleotides of one of the sequences of SEQ ID NOs 38-315 and a third primer having a sequence therein which is included within the sequence of said first primer; performing a first polymerase chain reaction with said first pair of nested primers to generate a first PCR product; contacting said first PCR product with a second pair of primers, said second pair of primers comprising a fourth primer, said fourth primer comprising at least 15 consecutive nucleotides of said sequence of one of SEQ ID NOs: 38-315, and a fifth primer, said fourth and fifth primers being capable of hybridizing to sequences within said first PCR product; and performing a second polymerase chain reaction, thereby generating a second PCR product.

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One aspect of the present invention is an isolated or purified cDNA encoding a human secretory protein, said human secretory protein comprising the protein encoded by one of SEQ ID NOs 38-315, or a fragment thereof of at least 10 amino acids, said cDNA being obtainable by the method of the preceding paragraph. In one embodiment, the cDNA comprises the full protein coding sequence partially included in one of the sequences of SEQ ID NOs: 38-315.

Another aspect of the present invention is the method described four paragraphs above in which the second cDNA strand is made by contacting said first cDNA strand with a second primer comprising at least 15 consecutive nucleotides of the sequences of SEQ ID NOs: 38-315; hybridizing said second primer to said first strand cDNA; and extending said hybridized second primer to generate said second cDNA strand.

Another aspect of the present invention is an isolated or purified cDNA encoding a human secretory protein, said human secretory protein comprising the protein partially encoded by one of SEQ ID NOs 38-315 or comprising a fragment thereof of at least 10 amino acids, said cDNA being obtainable by the method described in the preceding paragraph. In one embodiment, the cDNA comprises the full protein coding sequence partially included in of one of the sequences of SEQ ID NOs: 38-315.

Another aspect of the present invention is a method of making a protein comprising one of the sequences of SEQ ID NOs: 316-593, comprising the steps of obtaining a cDNA encoding the full protein sequence partially included in one of the sequences of sequence of SEQ ID NOs: 38-315; inserting said cDNA in an expression vector such that said cDNA is

operably linked to a promoter, introducing said expression vector into a host cell whereby said host cell produces the protein encoded by said cDNA; and isolating said protein.

Another aspect of the present invention is an isolated protein obtainable by the method described in the preceding paragraph.

Another aspect of the present invention is a method of obtaining a promoter DNA comprising the steps of obtaining DNAs located upstream of the nucleic acids of SEQ ID NOs: 38-315 or the sequences complementary thereto; screening said upstream DNAs to identify a promoter capable of directing transcription initiation; and isolating said DNA comprising said identified promoter. In one embodiment, the obtaining step comprises chromosome walking from said nucleic acids of SEQ ID NOs: 38-315 or sequences complementary thereto. In another embodiment, the screening step comprises inserting said upstream sequences into a promoter reporter vector. In another embodiment, the screening step comprises identifying motifs in said upstream DNAs which are transcription factor binding sites or transcription start sites.

Another aspect of the present invention is an isolated promoter obtainable by the method described above.

Another aspect of the present invention is an isolated or purified protein comprising one of the sequences of SEQ ID NOs: 316-593.

Another aspect of the present invention is the inclusion of at least one of the sequences of SEQ ID NOs: 38-315, or one of the sequences complementary to the sequences of SEQ ID NOs: 38-315, or a fragment thereof of at least 15 consecutive nucleotides in an array of discrete ESTs or fragments thereof of at least 15 nucleotides in length. In one embodiment, the array includes at least two of the sequences of SEQ ID NOs: 38-315, the sequences complementary to the sequences of SEQ ID NOs: 38-315, or fragments thereof of at least 15 consecutive nucleotides. In another embodiment, the array includes at least five of the sequences of SEQ ID NOs: 38-315, the sequences complementary to the sequences of SEQ ID NOs: 38-315, or fragments thereof of at least 15 consecutive nucleotides.

Another aspect of the present invention is a promoter having a sequence selected from the group consisting of SEQ ID NOs: 31, 34, and 37.

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Brief Description of the Drawings

Figure 1 is a summary of a procedure for obtaining cDNAs which have been selected to include the 5' ends of the mRNAs from which they derived.

Figure 2 shows the distribution of Von Heijne scores for 5' ESTs in each of the categories described herein and the probability that these 5' ESTs encode a signal peptide.

Figure 3 summarizes a general method used to clone and sequence extended cDNAs containing sequences adjacent to 5' ESTs.

Figure 4 (description of promoters structure isolated from SignalTag 5' ESTs) provides a schematic description of promoters isolated and the way they are assembled with the corresponding 5' tags.

Detailed Description of the Preferred Embodiment

Table IV is an analysis of the 43 amino acids located at the N terminus of all human SwissProt proteins to determine the frequency of false positives and false negatives using the techniques for signal peptide identification described herein.

Table V shows the distribution of 5' ESTs in each category described herein and the number of 5' ESTs in each category having a given minimum Von Heijne's score.

Table VI shows the distribution of 5' ESTs in each category described herein with respect to the tissue from which the 5' ESTs of the corresponding mRNA were obtained.

Table VII describes the transcription factor binding sites present in each of these promoters.

I. General Methods for Obtaining 5' ESTs derived from mRNAs with intact 5' ends

In order to obtain the 5' ESTs of the present invention, mRNAs with intact 5' ends must be obtained. Currently, there are two approaches for obtaining such mRNAs with intact 5' ends as described below: either chemical (1) or enzymatic (2).

1. Chemical Methods for Obtaining mRNAs having Intact 5' Ends

One of these approaches is a chemical modification method involving derivatization of the 5' ends of the mRNAs and selection of the derivatized mRNAs. The 5' ends of eukaryotic mRNAs possess a structure referred to as a "cap" which comprises a guanosine

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methylated at the 7 position. The cap is joined to the first transcribed base of the mRNA by a 5′, 5′-triphosphate bond. In some instances, the 5′ guanosine is methylated in both the 2 and 7 positions. Rarely, the 5′ guanosine is trimethylated at the 2, 7 and 7 positions. In the chemical method for obtaining mRNAs having intact 5′ ends, the 5′ cap is specifically derivatized and coupled to a reactive group on an immobilizing substrate. This specific derivatization is based on the fact that only the ribose linked to the methylated guanosine at the 5′ end of the mRNA and the ribose linked to the base at the 3′ terminus of the mRNA, possess 2′, 3′-cis diols.

Optionally, the 2', 3'-cis diol of the 3' terminal ribose may be chemically modified, substituted, converted, or eliminated, leaving only the ribose linked to the methylated guanosine at the 5' end of the mRNA with a 2', 3'-cis diol. A variety of techniques are available for eliminating the 2', 3'-cis diol on the 3' terminal ribose. For example, controlled alkaline hydrolysis may be used to generate mRNA fragments in which the 3' terminal ribose is a 3'-phosphate, 2'-phosphate or (2', 3')-cyclophosphate. Thereafter, the fragment which includes the original 3' ribose may be eliminated from the mixture through chromatography on an oligodT column. Alternatively, a base which lacks the 2', 3'-cis diol may be added to the 3' end of the mRNA using an RNA ligase such as T4 RNA ligase. Example 1 below describes a method for ligation of a nucleoside diphosphate to the 3' end of messenger RNA.

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EXAMPLE 1

Ligation of the Nucleoside Diphosphate pCp to the 3' End of mRNA.

One μg of RNA was incubated in a final reaction medium of 10 μl in the presence of 5 U of T_4 phage RNA ligase in the buffer provided by the manufacturer (Gibco-BRL), 40 U of the RNase inhibitor RNasin (Promega) and, 2 μl of ^{32}pCp (Amersham #PB 10208). The incubation was performed at 37°C for 2 hours or overnight at 7-8°C.

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Following modification or elimination of the 2', 3'-cis diol at the 3' ribose, the 2', 3'-cis diol present at the 5' end of the mRNA may be oxidized using reagents such as NaBH₃, NaBH₃CN, or sodium periodate, thereby converting the 2', 3'-cis diol to a dialdehyde.

Example 2 describes the oxidation of the 2', 3'-cis diol at the 5' end of the mRNA with sodium periodate.

EXAMPLE 2

Oxidation of 2', 3'-cis diol at the 5' End of the mRNA with Sodium Periodate

0.1 OD unit of either a capped oligoribonucleotide of 47 nucleotides (including the cap) or an uncapped oligoribonucleotide of 46 nucleotides were treated as follows. The oligoribonucleotides were produced by *in vitro* transcription using the transcription kit "AmpliScribe T7" (Epicentre Technologies). As indicated below, the DNA template for the RNA transcript contained a single cytosine. To synthesize the uncapped RNA, all four NTPs were included in the *in vitro* transcription reaction. To obtain the capped RNA, GTP was replaced by an analogue of the cap, m7G(5')ppp(5')G. This compound, recognized by the polymerase, was incorporated into the 5' end of the nascent transcript during the initiation of transcription but was not incorporated during the extension step. Consequently, the resulting RNA contained a cap at its 5' end. The sequences of the oligoribonucleotides produced by the *in vitro* transcription reaction were:

+Cap:

5'm7GpppGCAUCCUACUCCCAUCCAAUUCCACCCUAACUCCUCCCAUCUCCAC-3' (SEQ ID NO:1)

20 -Cap:

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5'-pppGCAUCCUACUCCAUCCAAUUCCACCUAACUCCUCCAUCUCCAC-3' (SEQ ID NO:2)

The oligoribonucleotides were dissolved in 9 μ l of acetate buffer (0.1 M sodium acetate, pH 5.2) and 3 μ l of freshly prepared 0.1 M sodium periodate solution. The mixture was incubated for 1 hour in the dark at 4°C or room temperature. Thereafter, the reaction was stopped by adding 4 μ l of 10% ethylene glycol. The product was ethanol precipitated, resuspended in at least 10 μ l of water or appropriate buffer and dialyzed against water.

The resulting aldehyde groups may then be coupled to molecules having a reactive amine group, such as hydrazine, carbazide, thiocarbazide or semicarbazide groups, in order to facilitate enrichment of the 5' ends of the mRNAs. Molecules having

reactive amine groups which are suitable for use in selecting mRNAs having intact 5' ends include avidin, proteins, antibodies, vitamins, ligands capable of specifically binding to receptor molecules, or oligonucleotides. Example 3 below describes the coupling of the resulting dialdehyde to biotin.

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EXAMPLE 3

Coupling of the Dialdehyde at the 5' End of Transcripts with Biotin

The oxidation product obtained in Example 2 was dissolved in 50 μ l of sodium acetate at a pH between 5 and 5.2 and 50 μ l of freshly prepared 0.02 M solution of biotin hydrazide in a methoxyethanol/water mixture (1:1) of formula:

In the compound used in these experiments, n=5. However, it will be appreciated that other commercially available hydrazides may also be used, such as molecules of the above formula in which n varies from 0 to 5. The mixture was then incubated for 2 hours at 37°C, precipitated with ethanol and dialyzed against distilled water. Example 4 demonstrates the specificity of the biotinylation reaction.

EXAMPLE 4

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Specificity of Biotinylation of Capped Transcripts

The specificity of the biotinylation for capped mRNAs was evaluated by gel electrophoresis of the following samples:

Sample 1. The 46 nucleotide uncapped *in vitro* transcript prepared as in Example 2 and labeled with ³²pCp as described in Example 1.

Sample 2. The 46 nucleotide uncapped *in vitro* transcript prepared as in Example 2, labeled with ³²pCp as described in Example 1, treated with the oxidation reaction of Example 2, and subjected to the biotinylation conditions of Example 3.

Sample 3. The 47 nucleotide capped *in vitro* transcript prepared as in Example 2 and labeled with ³²pCp as described in Example 1.

Sample 4. The 47 nucleotide capped *in vitro* transcript prepared as in Example 2, labeled with ³²pCp as described in Example 1, treated with the oxidation reaction of Example 2, and subjected to the biotinylation conditions of Example 3.

Samples 1 and 2 had identical migration rates, demonstrating that the uncapped RNAs were not oxidized and biotinylated. Sample 3 migrated more slowly than Samples 1 and 2, while Sample 4 exhibited the slowest migration. The difference in migration of the RNAs in Samples 3 and 4 demonstrates that the capped RNAs were specifically biotinylated.

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In some cases, mRNAs having intact 5' ends may be enriched by binding the molecule containing a reactive amine group to a suitable solid phase substrate such as the inside of the vessel containing the mRNAs, magnetic beads, chromatography matrices, or nylon or nitrocellulose membranes. For example, where the molecule having a reactive amine group is biotin, the solid phase substrate may be coupled to avidin or streptavidin. Alternatively, where the molecule having the reactive amine group is an antibody or receptor ligand, the solid phase substrate may be coupled to the cognate antigen or receptor. Finally, where the molecule having a reactive amine group comprises an oligonucleotide, the solid phase substrate may comprise a complementary oligonucleotide.

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The mRNAs having intact 5' ends may be released from the solid phase following the enrichment procedure. For example, where the dialdehyde is coupled to biotin hydrazide and the solid phase comprises streptavidin, the mRNAs may be released from the solid phase by simply heating to 95 degrees Celsius in 2% SDS. In some methods, the molecule having a reactive amine group may also be cleaved from the mRNAs having intact 5' ends following enrichment. Example 5 describes the capture of biotinylated mRNAs with streptavidin coated beads and the release of the biotinylated mRNAs from the beads following enrichment.

EXAMPLE 5

Capture and Release of Biotinylated mRNAs Using Streptavidin Coated Beads

The streptavidin coated magnetic beads were prepared according to the manufacturer's instructions (CPG Inc., USA). The biotinylated mRNAs were added to a

hybridization buffer (1.5 M NaCl, pH 5 - 6). After incubating for 30 minutes, the unbound and nonbiotinylated material was removed. The beads were then washed several times in water with 1% SDS. The beads thus obtained were incubated for 15 minutes at 95°C in water containing 2% SDS.

Example 6 demonstrates the efficiency with which biotinylated mRNAs were recovered from the streptavidin coated beads.

EXAMPLE 6

Efficiency of Recovery of Biotinylated mRNAs

The efficiency of the recovery procedure was evaluated as follows. Capped RNAs were labeled with ³²pCp, oxidized, biotinylated and bound to streptavidin coated beads as described above. Subsequently, the bound RNAs were incubated for 5, 15 or 30 minutes at 95°C in the presence of 2% SDS.

The products of the reaction were analyzed by electrophoresis on 12% polyacrylamide gels under denaturing conditions (7 M urea). The gels were subjected to autoradiography. During this manipulation, the hydrazone bonds were not reduced.

Increasing amounts of nucleic acids were recovered as incubation times in 2% SDS increased, demonstrating that biotinylated mRNAs were efficiently recovered.

In an alternative method for obtaining mRNAs having intact 5' ends, an oligonucleotide which has been derivatized to contain a reactive amine group is specifically coupled to mRNAs having an intact cap. Preferably, the 3' end of the mRNA is blocked prior to the step in which the aldehyde groups are joined to the derivatized oligonucleotide, as described above, so as to prevent the derivatized oligonucleotide from being joined to the 3' end of the mRNA using T4 RNA ligase as described in example 1. However, as discussed above, blocking the 3' end of the mRNA is an optional step. Derivatized oligonucleotides may be prepared as described in Example 7.

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EXAMPLE 7

Derivatization of Oligonucleotides

An oligonucleotide phosphorylated at its 3' end was converted to a 3' hydrazide in 3' by treatment with an aqueous solution of hydrazine or of dihydrazide of the formula $H_2N(R1)NH_2$ at about 1 to 3 M, and at pH 4.5 at a temperature of 8°C overnight. This incubation was performed in the presence of a carbodiimide type agent soluble in water such as 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide at a final concentration of 0.3 M.

The derivatized oligonucleotide was then separated from the other agents and products using a standard technique for isolating oligonucleotides.

As discussed above, the mRNAs to be enriched may be treated to eliminate the 3' OH groups which may be present thereon. This may be accomplished by enzymatic ligation of sequences lacking a 3' OH, such as pCp, as described in Example 1. Alternatively, the 3' OH groups may be eliminated by alkaline hydrolysis as described in Example 8 below.

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EXAMPLE 8

Elimination of 3' OH Groups of mRNA Using Alkaline Hydrolysis

In a total volume of $100 \,\mu l$ of $0.1 \,N$ sodium hydroxide, $1.5 \,\mu g$ mRNA is incubated for 40 to 60 minutes at 4°C. The solution is neutralized with acetic acid and precipitated with ethanol.

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Following the optional elimination of the 3' OH groups, the diol groups at the 5' ends of the mRNAs are oxidized as described below in Example 9.

EXAMPLE 9

Oxidation of Diols of mRNA

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Up to 1 OD unit of RNA was dissolved in 9 μ l of buffer (0.1 M sodium acetate, pH 6-7) or water and 3 μ l of freshly prepared 0.1 M sodium periodate solution. The reaction was incubated for 1 h in the dark at 4°C or room temperature. Following the incubation, the reaction was stopped by adding 4 μ l of 10% ethylene glycol. Thereafter the mixture was incubated at room temperature for 15 minutes. After ethanol precipitation, the product was resuspended in at least 10 μ l of water or appropriate buffer and dialyzed against water.

Following oxidation of the diol groups at the 5' ends of the mRNAs, the derivatized oligonucleotide was joined to the resulting aldehydes as described in Example 10.

EXAMPLE 10

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Ligature of Aldehydes of mRNA to Derivatized Oligonucleotides

The oxidized mRNA was dissolved in an acidic medium such as 50 µl of sodium acetate pH 4-6. Fifty µl of a solution of the derivatized oligonucleotide were added in order to obtain an mRNA:derivatized oligonucleotide ratio of 1:20. The mixture was reduced with a borohydride and incubated for 2 h at 37°C or overnight (14 h) at 10°C. The mixture was then ethanol precipitated, resuspended in 10 µl or more of water or appropriate buffer and dialyzed against distilled water. If desired, the resulting product may be analyzed using acrylamide gel electrophoresis, HPLC analysis, or other conventional techniques.

Following the attachment of the derivatized oligonucleotide to the mRNAs, a reverse transcription reaction may be performed as described in Example 11 below.

EXAMPLE 11

Reverse Transcription of mRNAs Ligatured to Derivatized Oligonucleotides

An oligodeoxyribonucleotide was derivatized as follows. Three OD units of an oligodeoxyribonucleotide of sequence 5'ATCAAGAATTCGCACGAGACCATTA3' (SEQ ID NO:3) having 5'-OH and 3'-P ends were dissolved in 70 µl of a 1.5 M hydroxybenzotriazole solution, pH 5.3, prepared in dimethylformamide/water (75:25) containing 2 µg of 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide. The mixture was incubated for 2 h 30 min at 22°C and then precipitated twice in LiClO₄/acetone. The pellet was resuspended in 200 µl of 0.25 M hydrazine and incubated at 8°C from 3 to 14 h. Following the hydrazine reaction, the mixture was precipitated twice in LiClO₄/acetone.

The messenger RNAs to be reverse transcribed were extracted from blocks of placenta having sides of 2 cm which had been stored at -80°C. The total RNA was extracted using conventional acidic phenol techniques. Oligo-dT chromatography was used to purify the mRNAs. The integrity of the mRNAs was checked by Northern-blotting.

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The diol groups on 7 µg of the placental mRNAs were oxidized as described above in Example 9. The derivatized oligonucleotide was joined to the mRNAs as described in Example 10 above except that the precipitation step was replaced by an exclusion chromatography step to remove derivatized oligodeoxyribonucleotides which were not joined to mRNAs. Exclusion chromatography was performed as follows:

Ten ml of Ultrogel AcA34 (BioSepra#230151) gel, a mix of agarose and acrylamide, were equilibrated in 50 ml of a solution of 10 mM Tris pH 8.0, 300 mM NaCl, 1 mM EDTA, and 0.05% SDS. The mixture was allowed to sediment. The supernatant was eliminated and the gel was resuspended in 50 ml of buffer. This procedure was repeated 2 or 3 times.

A glass bead (diameter 3 mm) was introduced into a 2 ml disposable pipette (length 25 cm). The pipette was filled with the gel suspension until the height of the gel stabilized at 1 cm from the top of the pipette. The column was then equilibrated with 20 ml of equilibration buffer (10 mM Tris HCl pH 7.4, 20 mM NaCl).

Ten μ l of the mRNA which had reacted with the derivatized oligonucleotide were mixed in 39 μ l of 10 mM urea and 2 μ l of blue-glycerol buffer, which had been prepared by dissolving 5 mg of bromophenol blue in 60% glycerol (v/v), and passing the mixture through a 0.45 μ m diameter filter.

The column was then loaded with the mRNAs coupled to the oligonucleotide. As soon as the sample had penetrated, equilibration buffer was added. Hundred µl fractions were then collected. Derivatized oligonucleotide which had not been attached to mRNA appeared in fraction 16 and later fractions. Thus, fractions 3 to 15 were combined and precipitated with ethanol.

To determine whether the derivatized oligonucleotide was actually linked to mRNA, one tenth of the combined fractions were spotted twice on a nylon membrane and hybridized to a radioactive probe using conventional techniques. The ³²P labeled probe used in these hybridizations was an oligodeoxyribonucleotide of sequence 5'TAATGGTCTCGTGCGAATTCTTGAT3' (SEQ ID NO:4) anticomplementary to the derivatized oligonucleotide. A signal observed after autoradiography, indicated that the derivatized oligonucleotide had been truly joined to the mRNA.

The remaining nine tenth of the mRNAs which had reacted with the derivatized oligonucleotide was reverse transcribed as follows. A reverse transcription reaction was

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carried out with reverse transcriptase following the manufacturer's instructions and 50 pmol of nonamers with random sequence as primers.

To ensure that reverse transcription had been carried out through the cap structure, two types of experiments were performed.

In the first approach, after elimination of RNA of the cDNA:RNA heteroduplexes obtained from the reverse transcription reaction by an alkaline hydrolysis, a portion of the resulting single stranded cDNAs was spotted on a positively charged membrane and hybridized, using conventional methods, to a ³²P labeled probe having a sequence identical to that of the derivatized oligonucleotide. Control spots containing, 1 pmol, 100 fmol, 50 fmol, 10 fmol and 1 fmol of a control oligodeoxyribonucleotide of sequence identical to that of the derivatized oligonucleotide were included. The signal observed in the spots containing the cDNA indicated that approximately 15 fmol of the derivatized oligonucleotide had been reverse transcribed. These results demonstrate that the reverse transcription can be performed through the cap and, in particular, that reverse transcriptase crosses the 5'-P-P-P-5' bond of the cap of eukaryotic messenger RNAs.

In the second type of experiment, the single stranded cDNAs obtained from the above first strand synthesis were used as template for PCR reactions. Two types of reactions were carried out. First, specific amplification of the mRNAs for alpha globin, dehydrogenase, pp15 and elongation factor E4 were carried out using the following pairs of oligodeoxyribonucleotide primers.

alpha-globin

GLO-S: 5'CCG ACA AGA CCA ACG TCA AGG CCG C3' (SEQ ID NO:5) GLO-As: 5'TCA CCA GCA GGC AGT GGC TTA GGA G 3' (SEQ ID NO:6)

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dehydrogenase

3 DH-S: 5'AGT GAT TCC TGC TAC TTT GGA TGG C3' (SEQ ID NO:7)
3 DH-As: 5'GCT TGG TCT TGT TCT GGA GTT TAG A3' (SEQ ID NO:8)

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pp15

PP15-S: 5'TCC AGA ATG GGA GAC AAG CCA ATT T3' (SEQ ID NO:9)

PP15-As: 5'AGG GAG GAG GAA ACA GCG TGA GTC C3' (SEQ ID NO:10)

Elongation factor E4

EFA1-S: 5'ATG GGA AAG GAA AAG ACT CAT ATC A3' (SEQ ID NO:11)

5 EF1A-As: 5'AGC AGC AAC AAT CAG GAC AGC ACA G3' (SEQ ID NO:12)

Second, non specific amplifications were also carried out with the antisense oligodeoxyribonucleotides of the pairs described above and with a primer derived from the sequence of the derivatized oligodeoxyribonucleotide (5'ATCAAGAATTCGCACGAGACCATTA3') (SEQ ID NO:13).

One twentieth of the following RT-PCR product samples were run on a 1.5% agarose gel and stained with ethidium bromide.

- Sample 1: The products of a PCR reaction using the globin primers of SEQ ID NOs 5 and 6 in the presence of cDNA.
- Sample 2: The products of a PCR reaction using the globin primers of SEQ ID NOs and 6 in the absence of added cDNA.
 - Sample 3: The products of a PCR reaction using the dehydrogenase primers of SEQ ID NOs 7 and 8 in the presence of cDNA.
- Sample 4: The products of a PCR reaction using the dehydrogenase primers of SEQ ID NOs 7 and 8 in the absence of added cDNA.
 - Sample 5: The products of a PCR reaction using the pp15 primers of SEQ ID NOs 9 and 10 in the presence of cDNA.
 - Sample 6: The products of a PCR reaction using the pp15 primers of SEQ ID NOs 9 and 10 in the absence of added cDNA.
- Sample 7: The products of a PCR reaction using the EIF4 primers of SEQ ID NOs 11 and 12 in the presence of added cDNA.
 - Sample 8: The products of a PCR reaction using the EIF4 primers of SEQ ID NOs 11 and 12 in the absence of added cDNA.
- A band of the size expected for the PCR product was observed only in samples 1, 3, 3 and 7, thus indicating the presence of the corresponding sequence in the cDNA population.

PCR reactions were also carried out with the antisense oligonucleotides of the globin and dehydrogenase primers (SEQ ID NOs 6 and 8) and an oligonucleotide whose sequence corresponds to that of the derivatized oligonucleotide. The presence of PCR products of the expected size in the samples equivalent to above samples 1 and 3 indicated that the derivatized oligonucleotide had been linked to mRNA.

The above examples summarize the chemical procedure for enriching mRNAs for those having intact 5' ends as illustrated in Figure 1. Further detail regarding the chemical approaches for obtaining such mRNAs are disclosed in International Application No. WO96/34981, published November 7, 1996, which is incorporated herein by reference. Strategies based on the above chemical modifications to the 5' cap structure may be utilized to generate cDNAs selected to include the 5' ends of the mRNAs from which they derived. In one version of such procedures, the 5' ends of the mRNAs are modified as described Thereafter, a reverse transcription reaction is conducted to extend a primer complementary to the 5' end of the mRNA. Single stranded RNAs are eliminated to obtain a population of cDNA/mRNA heteroduplexes in which the mRNA includes an intact 5' end. The resulting heteroduplexes may be captured on a solid phase coated with a molecule capable of interacting with the molecule used to derivatize the 5' end of the mRNA. Thereafter, the strands of the heteroduplexes are separated to recover single stranded first cDNA strands which include the 5' end of the mRNA. Second strand cDNA synthesis may then proceed using conventional techniques. For example, the procedures disclosed in WO 96/34981 or in Carninci. et al., Genomics 37:327-336, 1996, the disclosures of which are incorporated herein by reference, may be employed to select cDNAs which include the sequence derived from the 5' end of the coding sequence of the mRNA.

Following ligation of the oligonucleotide tag to the 5' cap of the mRNA, a reverse transcription reaction is conducted to extend a primer complementary to the mRNA to the 5' end of the mRNA. Following elimination of the RNA component of the resulting heteroduplex using standard techniques, second strand cDNA synthesis is conducted with a primer complementary to the oligonucleotide tag.

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2. Enzymatic Methods for Obtaining mRNAs having Intact 5' Ends

Other techniques for selecting cDNAs extending to the 5' end of the mRNA from which they are derived are fully enzymatic. Some versions of these techniques are disclosed in Dumas Milne Edwards J.B. (Doctoral Thesis of Paris VI University, Le clonage des ADNc complets: difficultes et perspectives nouvelles. Apports pour l'etude de la regulation de l'expression de la tryptophane hydroxylase de rat, 20 Dec. 1993), EP0 625572 and Kato et al., Gene 150:243-250, 1994, the disclosures of which are incorporated herein by reference.

Briefly, in such approaches, isolated mRNA is treated with alkaline phosphatase to remove the phosphate groups present on the 5' ends of uncapped incomplete mRNAs. Following this procedure, the cap present on full length mRNAs is enzymatically removed with a decapping enzyme such as T4 polynucleotide kinase or tobacco acid pyrophosphatase. An oligonucleotide, which may be either a DNA oligonucleotide or a DNA-RNA hybrid oligonucleotide having RNA at its 3' end, is then ligated to the phosphate present at the 5' end of the decapped mRNA using T4 RNA ligase. The oligonucleotide may include a restriction site to facilitate cloning of the cDNAs following their synthesis. Example 12 below describes one enzymatic method based on the doctoral thesis of Dumas.

EXAMPLE 12

Enzymatic Approach for Obtaining 5' ESTs

Twenty micrograms of PolyA+ RNA were dephosphorylated using Calf Intestinal Phosphatase (Biolabs). After a phenol chloroform extraction, the cap structure of mRNA was hydrolysed using the Tobacco Acid Pyrophosphatase (purified as described by Shinshi *et al.*., *Biochemistry* 15: 2185-2190, 1976) and a hemi 5'DNA/RNA-3' oligonucleotide having an unphosphorylated 5' end, a stretch of adenosine ribophosphate at the 3' end, and an EcoRI site near the 5' end was ligated to the 5'P ends of mRNA using the T4 RNA ligase (Biolabs). Oligonucleotides suitable for use in this procedure are preferably 30 to 50 bases in length. Oligonucleotides having an unphosphorylated 5' end may be synthesized by adding a fluorochrome at the 5' end. The inclusion of a stretch of adenosine ribophosphates at the 3' end of the oligonucleotide increases ligation efficiency. It will be appreciated that the oligonucleotide may contain cloning sites other than EcoRI.

Following ligation of the oligonucleotide to the phosphate present at the 5' end of the decapped mRNA, first and second strand cDNA synthesis is carried out using conventional methods or those specified in EP0 625,572 and Kato et al. supra, and Dumas Milne Edwards, supra, the disclosures of which are incorporated herein by reference. The resulting cDNA may then be ligated into vectors such as those disclosed in Kato et al., supra or other nucleic acid vectors known to those skilled in the art using techniques such as those described in Sambrook et al., Molecular Cloning: A Laboratory Manual 2d Ed., Cold Spring Harbor Laboratory Press, 1989, the disclosure of which is incorporated herein by reference.

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II. Obtention and Characterization of the 5' ESTs of the Present Invention

The 5' ESTs of the present invention were obtained using the aforementioned chemical and enzymatic approaches for enriching mRNAs for those having intact 5' ends as decribed below.

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1. Obtention of 5' ESTS Using mRNAs with Intact 5' Ends

First, mRNAs were prepared as described in Example 13 below.

EXAMPLE 13

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Preparation of mRNA With Intact 5' Ends

Total human RNAs or polyA⁺ RNAs derived from 29 different tissues were respectively purchased from LABIMO and CLONTECH and used to generate 44 cDNA libraries as follows. The purchased RNA had been isolated from cells or tissues using acid guanidium thiocyanate-phenol-chloroform extraction (Chomczyniski and Sacchi, *Analytical Biochemistry* 162:156-159, 1987). PolyA⁺ RNA was isolated from total RNA (LABIMO) by two passes of oligo dT chromatography, as described by Aviv and Leder, *Proc. Natl. Acad. Sci. USA* 69:1408-1412, 1972 in order to eliminate ribosomal RNA.

The quality and the integrity of the polyA+ RNAs were checked. Northern blots hybridized with a globin probe were used to confirm that the mRNAs were not degraded. Contamination of the polyA+ mRNAs by ribosomal sequences was checked using Northern blots and a probe derived from the sequence of the 28S rRNA. Preparations of mRNAs with

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less than 5% of rRNAs were used in library construction. To avoid constructing libraries with RNAs contaminated by exogenous sequences (prokaryotic or fungal), the presence of bacterial 16S ribosomal sequences or of two highly expressed fungal mRNAs was examined using PCR.

Following preparation of the mRNAs, the above described chemical and/or the enzymatic procedures for enriching mRNAs for thoses having intact 5' ends were employed to obtain 5' ESTs from various tissues. In both approaches, an oligonucleotide tag was attached to the 5' ends of the mRNAs. The oligonucleotide tag had an EcoRI site therein to facilitate later cloning procedures. To facilitate the processing of single stranded and double stranded cDNA obtained in the construction of the librairies, the same nucleotidic sequence was used to design the ligated oligonucleotide in both chemical and enzymatic approaches. Nevertheless, in the chemical procedure, the tag used was an oligodeoxyribonucleotide which was linked to the cap of the mRNA whereas in the enzymatic ligation, the tag was a chimeric hemi 5'DNA/RNA3' oligonucleotide which was ligated to the 5' end of decapped mRNA as described in example 12.

Following attachment of the oligonucleotide tag to the mRNA by either the chemical or enzymatic methods, the integrity of the mRNA was examined by performing a Northern blot with 200 to 500 ng of mRNA using a probe complementary to the oligonucleotide tag before performing the first strand synthesis as described in example 14.

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EXAMPLE 14

cDNA Synthesis Using mRNA Templates Having Intact 5' Ends

For the mRNAs joined to oligonucleotide tags using both the chemical and enzymatic methods, first strand cDNA synthesis was performed using the Superscript II (Gibco BRL) or the Rnase H Minus M-MLV (Promega) reverse transcriptase with random nonamers as primers. In order to protect internal EcoRI sites in the cDNA from digestion at later steps in the procedure, methylated dCTP was used for first strand synthesis. After removal of RNA by an alkaline hydrolysis, the first strand of cDNA was precipitated using isopropanol in order to eliminate residual primers.

For both the chemical and the enzymatic methods, the second strand of the cDNA was synthesized with a Klenow fragment using a primer corresponding to the 5' end of the

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ligated oligonucleotide described in Example 12. Preferably, the primer is 20-25 bases in length. Methylated dCTP was also used for second strand synthesis in order to protect internal EcoRI sites in the cDNA from digestion during the cloning process.

Following cDNA synthesis, the cDNAs were cloned into pBlueScript as described in Example 15 below.

EXAMPLE 15

Cloning of cDNAsderived from mRNA with intact 5' ends into BlueScript

Following second strand synthesis, the ends of the cDNA were blunted with T4 DNA polymerase (Biolabs) and the cDNA was digested with EcoRI. Since methylated dCTP was used during cDNA synthesis, the EcoRI site present in the tag was the only hemi-methylated site, hence the only site susceptible to EcoRI digestion. The cDNA was then size fractionated using exclusion chromatography (AcA, Biosepra) and fractions corresponding to cDNAs of more than 150 bp were pooled and ethanol precipitated. The cDNA was directionally cloned into the SmaI and EcoRI ends of the phagemid pBlueScript vector (Stratagene). The ligation mixture was electroporated into bacteria and propagated under appropriate antibiotic selection.

Clones containing the oligonucleotide tag attached were then selected as described in Example 16 below.

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EXAMPLE 16

Selection of Clones Having the Oligonucleotide Tag Attached Thereto

The plasmid DNAs containing 5' EST libraries made as described above were purified (Qiagen). A positive selection of the tagged clones was performed as follows. Briefly, in this selection procedure, the plasmid DNA was converted to single stranded DNA using gene II endonuclease of the phage F1 in combination with an exonuclease (Chang et al., Gene 127:95-8, 1993) such as exonuclease III or T7 gene 6 exonuclease. The resulting single stranded DNA was then purified using paramagnetic beads as described by Fry et al., Biotechniques, 13: 124-131, 1992. In this procedure, the single stranded DNA was hybridized with a biotinylated oligonucleotide having a sequence corresponding to the 3' end of the oligonucleotide described in Example 13. Preferably, the primer has a length of 20-25

bases. Clones including a sequence complementary to the biotinylated oligonucleotide were captured by incubation with streptavidin coated magnetic beads followed by magnetic selection. After capture of the positive clones, the plasmid DNA was released from the magnetic beads and converted into double stranded DNA using a DNA polymerase such as the ThermoSequenase obtained from Amersham Pharmacia Biotech. Alternatively, protocoles such as the one described in the Gene Trapper kit available from Gibco BRL may be used. The double stranded DNA was then electroporated into bacteria. The percentage of positive clones having the 5' tag oligonucleotide was estimated to typically rank between 90 and 98% using dot blot analysis.

Following electroporation, the libraries were ordered in 384-microtiter plates (MTP). A copy of the MTP was stored for future needs. Then the libraries were transferred into 96 MTP and sequenced as described below.

EXAMPLE 17

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Sequencing of Inserts in Selected Clones

Plasmid inserts were first amplified by PCR on PE 9600 thermocyclers (Perkin-Elmer, Applied Biosystems Division, Foster City, CA), using standard SETA-A and SETA-B primers (Genset SA), AmpliTaqGold (Perkin-Elmer), dNTPs (Boehringer), buffer and cycling conditions as recommended by the Perkin-Elmer Corporation.

PCR products were then sequenced using automatic ABI Prism 377 sequencers (Perkin Elmer). Sequencing reactions were performed using PE 9600 thermocyclers with standard dye-primer chemistry and ThermoSequenase (Amersham Pharmacia Biotech). The primers used were either T7 or 21M13 (available from Genset SA) as appropriate. The primers were labeled with the JOE, FAM, ROX and TAMRA dyes. The dNTPs and ddNTPs used in the sequencing reactions were purchased from Boehringer. Sequencing buffer, reagent concentrations and cycling conditions were as recommended by Amersham.

Following the sequencing reaction, the samples were precipitated with ethanol, resuspended in formamide loading buffer, and loaded on a standard 4% acrylamide gel. Electrophoresis was performed for 2.5 hours at 3000V on an ABI 377 sequencer, and the sequence data were collected and analyzed using the ABI Prism DNA Sequencing Analysis Software, version 2.1.2.

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2. Computer analysis of the Obtained 5' ESTs: Construction of NetGene and SignalTag databases

The sequence data from the 44 cDNA libraries made as described above were transferred to a proprietary database, where quality control and validation steps were performed. A proprietary base-caller, working using a Unix system, automatically flagged suspect peaks, taking into account the shape of the peaks, the inter-peak resolution, and the noise level. The proprietary base-caller also performed an automatic trimming. Any stretch of 25 or fewer bases having more than 4 suspect peaks was considered unreliable and was discarded. Sequences corresponding to cloning vector or ligation oligonucleotides were automatically removed from the EST sequences. However, the resulting EST sequences may contain 1 to 5 bases belonging to the above mentioned sequences at their 5' end. If needed, these can easily be removed on a case to case basis.

Following sequencing as described above, the sequences of the 5' ESTs were entered in NetGeneTM, a proprietary database called for storage and manipulation as described below. It will be appreciated by those skilled in the art that the data could be stored and manipulated on any medium which can be read and accessed by a computer. Computer readable media include magnetically, optically, or electronically readable media. For example, the computer readable media may be a hard disc, a floppy disc, a magnetic tape, CD-ROM, RAM, or ROM as well as other types of other media known to those skilled in the art.

In addition, the sequence data may be stored and manipulated in a variety of data processor programs in a diversity of formats. For instance, the sequence data may be stored as text in a word processing file, such as Microsoft WORD or WORDPERFECT or as an ASCII file in a variety of database programs familiar to those of skill in the art, such as DB2, SYBASE, or ORACLE.

The computer readable media on which the sequence information is stored may be in a personal computer, a network, a server or other computer systems known to those skilled in the art. The computer or other system preferably includes the storage media described above, and a processor for accessing and manipulating the sequence data. Once the sequence data has been stored, it may be manipulated and searched to locate those stored sequences which contain a desired nucleic acid sequence or which encode a protein having a particular functional domain. For example, the stored sequence information may be compared to other

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known sequences to identify homologies, motifs implicated in biological function, or structural motifs.

Programs which may be used to search or compare the stored sequences include the MacPattern (EMBL), BLAST, and BLAST2 program series (NCBI), basic local alignment search tool programs for nucleotide (BLASTN) and peptide (BLASTX) comparisons (Altschul et al, J. Mol. Biol. 215: 403, 1990) and FASTA (Pearson and Lipman, Proc. Natl. Acad. Sci. USA 85: 2444, 1988). The BLAST programs then extend the alignments on the basis of defined match and mismatch criteria.

Motifs which may be detected using the above programs and those described in Example 28 include sequences encoding leucine zippers, helix-turn-helix motifs, glycosylation sites, ubiquitination sites, alpha helices, and beta sheets, signal sequences encoding signal peptides which direct the secretion of the encoded proteins, sequences implicated in transcription regulation such as homeoboxes, acidic stretches, enzymatic active sites, substrate binding sites, and enzymatic cleavage sites.

Before searching the cDNAs in the NetGene[™] database for sequence motifs of interest, cDNAs derived from mRNAs which were not of interest were identified and eliminated from further consideration as described in Example 18 below.

EXAMPLE 18

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Elimination of Undesired Sequences from Further Consideration

5' ESTs in the NetGene™ database which were derived from undesired sequences such as transfer RNAs, ribosomal RNAs, mitochondrial RNAs, prokaryotic RNAs, fungal RNAs, Alu sequences, L1 sequences, or repeat sequences were identified using the FASTA and BLASTN programs with the parameters listed in Table I.

To eliminate 5' ESTs encoding tRNAs from further consideration, the 5' EST sequences were compared to the sequences of 1190 known tRNAs obtained from EMBL release 38, of which 100 were human. The comparison was performed using FASTA on both strands of the 5' ESTs. Sequences having more than 80% homology over more than 60 nucleotides were identified as tRNA. Of the 144,341 sequences screened, 26 were identified as tRNAs and eliminated from further consideration.

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To eliminate 5' ESTs encoding rRNAs from further consideration, the 5' EST sequences were compared to the sequences of 2497 known rRNAs obtained from EMBL release 38, of which 73 were human. The comparison was performed using BLASTN on both strands of the 5' ESTs with the parameter S=108. Sequences having more than 80% homology over stretches longer than 40 nucleotides were identified as rRNAs. Of the 144,341 sequences screened, 3,312 were identified as rRNAs and eliminated from further consideration.

To eliminate 5' ESTs encoding mtRNAs from further consideration, the 5' EST sequences were compared to the sequences of the two known mitochondrial genomes for which the entire genomic sequences are available and all sequences transcribed from these mitochondrial genomes including tRNAs, rRNAs, and mRNAs for a total of 38 sequences. The comparison was performed using BLASTN on both strands of the 5' ESTs with the parameter S=108. Sequences having more than 80% homology over stretches longer than 40 nucleotides were identified as mtRNAs. Of the 144,341 sequences screened, 6,110 were identified as mtRNAs and eliminated from further consideration.

Sequences which might have resulted from exogenous contaminants were eliminated from further consideration by comparing the 5' EST sequences to release 46 of the EMBL bacterial and fungal divisions using BLASTN with the parameter S=144. All sequences having more than 90% homology over at least 40 nucleotides were identified as exogenous contaminants. Of the 42 cDNA libraries examined, the average percentages of prokaryotic and fungal sequences contained therein were 0.2% and 0.5% respectively. Among these sequences, only one could be identified as a sequence specific to fungi. The others were either fungal or prokaryotic sequences having homologies with vertebrate sequences or including repeat sequences which had not been masked during the electronic comparison.

In addition, the 5' ESTs were compared to 6093 Alu sequences and 1115 L1 sequences to mask 5' ESTs containing such repeat sequences. 5' ESTs including THE and MER repeats, SSTR sequences or satellite, micro-satellite, or telomeric repeats were also eliminated from further consideration. On average, 11.5% of the sequences in the libraries contained repeat sequences. Of this 11.5%, 7% contained Alu repeats, 3.3% contained L1 repeats and the remaining 1.2% were derived from the other screened types of repetitive sequences. These percentages are consistent with those found in cDNA libraries prepared by

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other groups. For example, the cDNA libraries of Adams *et al.* contained between 0% and 7.4% Alu repeats depending on the source of the RNA which was used to prepare the cDNA library (Adams *et al.*, *Nature* 377:174, 1996).

The sequences of those 5' ESTs remaining after the elimination of undesirable sequences were compared with the sequences of known human mRNAs to determine the accuracy of the sequencing procedures described above.

EXAMPLE 19

Measurement of Sequencing Accuracy by Comparison to Known Sequences

To further determine the accuracy of the sequencing procedure described above, the sequences of 5' ESTs derived from known sequences were identified and compared to the original known sequences. First, a FASTA analysis with overhangs shorter than 5 bp on both ends was conducted on the 5' ESTs to identify those matching an entry in the public human mRNA database. The 6655 5' ESTs which matched a known human mRNA were then realigned with their cognate mRNA and dynamic programming was used to include substitutions, insertions, and deletions in the list of "errors" which would be recognized. Errors occurring in the last 10 bases of the 5' EST sequences were ignored to avoid the inclusion of spurious cloning sites in the analysis of sequencing accuracy.

This analysis revealed that the sequences incorporated in the NetGene[™] database had an accuracy of more than 99.5%.

To determine the efficiency with which the above selection procedures select cDNAs which include the 5' ends of their corresponding mRNAs, the following analysis was performed.

EXAMPLE 20

Determination of Efficiency of 5' EST Selection

To determine the efficiency at which the above selection procedures isolated 5' ESTs which included sequences close to the 5' end of the mRNAs from which they derived, the sequences of the ends of the 5' ESTs derived from the elongation factor 1 subunit α and

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ferritin heavy chain genes were compared to the known cDNA sequences of these genes. Since the transcription start sites of both genes are well characterized, they may be used to determine the percentage of derived 5' ESTs which included the authentic transcription start sites.

For both genes, more than 95% of the obtained 5' ESTs actually included sequences close to or upstream of the 5' end of the corresponding mRNAs.

To extend the analysis of the reliability of the procedures for isolating 5' ESTs from ESTs in the NetGeneTM database, a similar analysis was conducted using a database composed of human mRNA sequences extracted from GenBank database release 97 for comparison. The 5' ends of more than 85% of 5' ESTs derived from mRNAs included in the GeneBank database were located close to the 5' ends of the known sequence. As some of the mRNA sequences available in the GenBank database are deduced from genomic sequences, a 5' end matching with these sequences will be counted as an internal match. Thus, the method used here underestimates the yield of ESTs including the authentic 5' ends of their corresponding mRNAs.

The EST libraries made above included multiple 5' ESTs derived from the same mRNA. The sequences of such 5' ESTs were compared to one another and the longest 5' ESTs for each mRNA were identified. Overlapping cDNAs were assembled into continuous sequences (contigs). The resulting continuous sequences were then compared to public databases to gauge their similarity to known sequences, as described in Example 21 below.

EXAMPLE 21

Clustering of the 5' ESTs and Calculation of Novelty Indices for cDNA Libraries

For each sequenced EST library, the sequences were clustered by the 5' end. Each sequence in the library was compared to the others with BLASTN2 (direct strand, parameters S=107). ESTs with High Scoring Segment Pairs (HSPs) at least 25 bp long, having 95% identical bases and beginning closer than 10 bp from each EST 5' end were grouped. The longest sequence found in the cluster was used as representative of the group. A global clustering between libraries was then performed leading to the definition of super-contigs.

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To assess the yield of new sequences within the EST libraries, a novelty rate (NR) was defined as: NR= 100 X (Number of new unique sequences found in the library/Total number of sequences from the library). Typically, novelty rating ranged between 10% and 41% depending on the tissue from which the EST library was obtained. For most of the libraries, the random sequencing of 5' EST libraries was pursued until the novelty rate reached 20%.

Following characterization as described above, the collection of 5' ESTs in NetGeneTM was screened to identify those 5' ESTs bearing potential signal sequences as described in Example 22 below.

EXAMPLE 22

Identification of Potential Signal Sequences in 5' ESTs

The 5' ESTs in the NetGeneTM database were screened to identify those having an uninterrupted open reading frame (ORF) longer than 45 nucleotides beginning with an ATG codon and extending to the end of the EST. Approximately half of the cDNA sequences in NetGeneTM contained such an ORF. The ORFs of these 5' ESTs were then searched to identify potential signal motifs using slight modifications of the procedures disclosed in Von Heijne, *Nucleic Acids Res.* 14:4683-4690, 1986, the disclosure of which is incorporated herein by reference. Those 5' EST sequences encoding a stretch of at least 15 amino acid long with a score of at least 3.5 in the Von Heijne signal peptide identification matrix were considered to possess a signal sequence. Those 5' ESTs which matched a known human mRNA or EST sequence and had a 5' end more than 20 nucleotides downstream of the known 5' end were excluded from further analysis. The remaining cDNAs having signal sequences therein were included in a database called SignalTagTM.

To confirm the accuracy of the above method for identifying signal sequences, the analysis of Example 23 was performed.

30 EXAMPLE 23

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The accuracy of the above procedure for identifying signal sequences encoding signal peptides was evaluated by applying the method to the 43 amino acids located at the N terminus of all human SwissProt proteins. The computed Von Heijne score for each protein was compared with the known characterization of the protein as being a secreted protein or a non-secreted protein. In this manner, the number of non-secreted proteins having a score higher than 3.5 (false positives) and the number of secreted proteins having a score lower than 3.5 (false negatives) could be calculated.

Using the results of the above analysis, the probability that a peptide encoded by the 5' region of the mRNA is in fact a genuine signal peptide based on its Von Heijne's score was calculated based on either the assumption that 10% of human proteins are secreted or the assumption that 20% of human proteins are secreted. The results of this analysis are shown in Figure 2 and in table IV.

Using the above method of identification of secretory proteins, 5' ESTs of the following polypeptides known to be secreted were obtained: human glucagon, gamma interferon induced monokine precursor, secreted cyclophilin-like protein, human pleiotropin, and human biotinidase precursor. Thus, the above method successfully identified those 5' ESTs which encode a signal peptide.

To confirm that the signal peptide encoded by the 5' ESTs actually functions as a signal peptide, the signal sequences from the 5' ESTs may be cloned into a vector designed for the identification of signal peptides. Such vectors are designed to confer the ability to grow in selective medium only to host cells containing a vector with an operably linked signal sequence. For example, to confirm that a 5' EST encodes a genuine signal peptide, the signal sequence of the 5' EST may be inserted upstream and in frame with a non-secreted form of the yeast invertase gene in signal peptide selection vectors such as those described in U.S. Patent No. 5,536,637, the disclosure of which is incorporated herein by reference. Growth of host cells containing signal sequence selection vectors with the correctly inserted 5' EST signal sequence confirms that the 5' EST encodes a genuine signal peptide.

Alternatively, the presence of a signal peptide may be confirmed by cloning the extended cDNAs obtained using the ESTs into expression vectors such as pXT1 (as described below in example 30), or by constructing promoter-signal sequence-reporter gene

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vectors which encode fusion proteins between the signal peptide and an assayable reporter protein. After introduction of these vectors into a suitable host cell, such as COS cells or NIH 3T3 cells, the growth medium may be harvested and analyzed for the presence of the secreted protein. The medium from these cells is compared to the medium from control cells containing vectors lacking the signal sequence or extended cDNA insert to identify vectors which encode a functional signal peptide or an authentic secreted protein.

Those 5' ESTs which encoded a signal peptide, as determined by the method of Example 22 above, were further grouped into four categories based on their homology to known sequences as described in Example 24 below.

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EXAMPLE 24

Categorization of 5' ESTs Encoding a Signal Peptide

Those 5' ESTs having a sequence not matching any known vertebrate sequence nor any publicly available EST sequence were designated "new." Of the sequences in the SignalTag™ database, 947 of the 5' ESTs having a Von Heijne's score of at least 3.5 fell into this category.

Those 5' ESTs having a sequence not matching any vertebrate sequence but matching a publicly known EST were designated "EST-ext", provided that the known EST sequence was extended by at least 40 nucleotides in the 5' direction. Of the sequences in the ṢignalTag™ database, 150 of the 5' ESTs having a Von Heijne's score of at least 3.5 fell into this category.

Those ESTs not matching any vertebrate sequence but matching a publicly known EST without extending the known EST by at least 40 nucleotides in the 5' direction were designated "EST." Of the sequences in the SignalTag[™] database, 599 of the 5' ESTs having a Von Heijne's score of at least 3.5 fell into this category.

Those 5' ESTs matching a human mRNA sequence but extending the known sequence by at least 40 nucleotides in the 5' direction were designated "VERT-ext." Of the sequences in the SignalTag™ database, 23 of the 5' ESTs having a Von Heijne's score of at least 3.5 fell into this category. Included in this category was a 5' EST which extended the known sequence of the human translocase mRNA by more than 200 bases in the 5' direction.

A 5' EST which extended the sequence of a human tumor suppressor gene in the 5' direction was also identified.

Table V shows the distribution of 5' ESTs in each category and the number of 5' ESTs in each category having a given minimum von Heijne's score.

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3. Evaluation of Spatial and Temporal Expression of mRNAs Corresponding to the 5'ESTs or Extended cDNAs

Each of the 5' ESTs was also categorized based on the tissue from which its corresponding mRNA was obtained, as described below in Example 25.

EXAMPLE 25

Categorization of Expression Patterns

Table VI shows the distribution of 5' ESTs in each of the above defined category with respect to the tissue from which the 5'ESTs of the corresponding mRNA were obtained.

Table II provides the sequence identification numbers of 5' EST sequences derived from prostate, the categories in which these sequences fall, and the von Heijne's score of the signal peptides which they encode. The 5' EST sequences and the amino acid sequences they encode are provided in the appended sequence listings. Table III provides the sequence ID numbers of the 5' ESTs and the sequences of the signal peptides which they encode. The sequences of the 5' ESTs and the polypeptides they encode are provided in the sequence listing appended hereto.

The sequences of DNA SEQ ID NOs: 38-315 can readily be screened for any errors therein and any sequence ambiguities can be resolved by resequencing a fragment containing such errors or ambiguities on both strands. Such fragments may be obtained from the plasmids stored in the inventors' laboratory or can be isolated using the techniques described herein. Resolution of any such ambiguities or errors may be facilitated by using primers which hybridize to sequences located close to the ambiguous or erroneous sequences. For example, the primers may hybridize to sequences within 50-75 bases of the ambiguity or error. Upon resolution of an error or ambiguity, the corresponding corrections can be made in the protein sequences encoded by the DNA containing the error or ambiguity.

In addition to categorizing the 5' ESTs with respect to their tissue of origin, the spatial and temporal expression patterns of the mRNAs corresponding to the 5' ESTs, as well as their expression levels, may be determined as described in Example 26 below. Characterization of the spatial and temporal expression patterns and expression levels of these mRNAs is useful for constructing expression vectors capable of producing a desired level of gene product in a desired spatial or temporal manner, as will be discussed in more detail below.

Furthermore, 5' ESTs whose corresponding mRNAs are associated with disease states may also be identified. For example, a particular disease may result from the lack of expression, over expression, or under expression of an mRNA corresponding to a 5' EST. By comparing mRNA expression patterns and quantities in samples taken from healthy individuals with those from individuals suffering from a particular disease, 5' ESTs responsible for the disease may be identified.

It will be appreciated that the results of the above characterization procedures for 5' ESTs also apply to extended cDNAs (obtainable as described below) which contain sequences adjacent to the 5' ESTs. It will also be appreciated that if desired, characterization may be delayed until extended cDNAs have been obtained rather than characterizing the ESTs themselves.

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EXAMPLE 26

Evaluation of Expression Levels and Patterns of mRNAs Corresponding to 5' ESTs or Extended cDNAs

Expression levels and patterns of mRNAs corresponding to 5' ESTs or extended cDNAs (obtainable as described below in example 27) may be analyzed by solution hybridization with long probes as described in International Patent Application No. WO 97/05277, the entire contents of which are hereby incorporated by reference. Briefly, a 5' EST, extended cDNA, or fragment thereof corresponding to the gene encoding the mRNA to be characterized is inserted at a cloning site immediately downstream of a bacteriophage (T3, T7 or SP6) RNA polymerase promoter to produce antisense RNA. Preferably, the 5' EST or extended cDNA has 100 or more nucleotides. The plasmid is linearized and transcribed in the

presence of ribonucleotides comprising modified ribonucleotides (*i.e.* biotin-UTP and DIG-UTP). An excess of this doubly labeled RNA is hybridized in solution with mRNA isolated from cells or tissues of interest. The hybridizations are performed under standard stringent conditions (40-50°C for 16 hours in an 80% formamide, 0.4 M NaCl buffer, pH 7-8). The unhybridized probe is removed by digestion with ribonucleases specific for single-stranded RNA (*i.e.* RNases CL3, T1, Phy M, U2 or A). The presence of the biotin-UTP modification enables capture of the hybrid on a microtitration plate coated with streptavidin. The presence of the DIG modification enables the hybrid to be detected and quantified by ELISA using an anti-DIG antibody coupled to alkaline phosphatase.

The 5' ESTs, extended cDNAs, or fragments thereof may also be tagged with nucleotide sequences for the serial analysis of gene expression (SAGE) as disclosed in UK Patent Application No. 2 305 241 A, the entire contents of which are incorporated by reference. In this method, cDNAs are prepared from a cell, tissue, organism or other source of nucleic acid for which gene expression patterns must be determined. The resulting cDNAs are separated into two pools. The cDNAs in each pool are cleaved with a first restriction endonuclease, called an anchoring enzyme, having a recognition site which is likely to be present at least once in most cDNAs. The fragments which contain the 5' or 3' most region of the cleaved cDNA are isolated by binding to a capture medium such as streptavidin coated beads. A first oligonucleotide linker having a first sequence for hybridization of an amplification primer and an internal restriction site for a so-called tagging endonuclease is ligated to the digested cDNAs in the first pool. Digestion with the second endonuclease produces short tag fragments from the cDNAs.

A second oligonucleotide having a second sequence for hybridization of an amplification primer and an internal restriction site is ligated to the digested cDNAs in the second pool. The cDNA fragments in the second pool are also digested with the tagging endonuclease to generate short tag fragments derived from the cDNAs in the second pool. The tags resulting from digestion of the first and second pools with the anchoring enzyme and the tagging endonuclease are ligated to one another to produce so-called ditags. In some embodiments, the ditags are concatamerized to produce ligation products containing from 2 to 200 ditags. The tag sequences are then determined and compared to the sequences of the 5' ESTs or extended cDNAs to determine which 5' ESTs or extended cDNAs are expressed

in the cell, tissue, organism, or other source of nucleic acids from which the tags were derived. In this way, the expression pattern of the 5' ESTs or extended cDNAs in the cell, tissue, organism, or other source of nucleic acids is obtained.

Quantitative analysis of gene expression may also be performed using arrays. As used herein, the term array means a one dimensional, two dimensional, or multidimensional arrangement of full length cDNAs (*i.e.* extended cDNAs which include the coding sequence for the signal peptide, the coding sequence for the mature protein, and a stop codon), extended cDNAs, 5' ESTs or fragments thereof of sufficient length to permit specific detection of gene expression. Preferably, the fragments are at least 15 nucleotides in length. More preferably, the fragments are at least 100 nucleotide long. More preferably, the fragments are more than 100 nucleotides in length. In some embodiments, the fragments may be more than 500 nucleotide long.

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For example, quantitative analysis of gene expression may be performed with full length cDNAs as defined below, extended cDNAs, 5' ESTs, or fragments thereof in a complementary DNA microarray as described by Schena *et al.* (*Science* 270:467-470, 1995; *Proc. Natl. Acad. Sci. U.S.A.* 93:10614-10619, 1996). Full length cDNAs, extended cDNAs, 5' ESTs or fragments thereof are amplified by PCR and arrayed from 96-well microtiter plates onto silylated microscope slides using high-speed robotics. Printed arrays are incubated in a humid chamber to allow rehydration of the array elements and rinsed, once in 0.2% SDS for 1 min, twice in water for 1 min and once for 5 min in sodium borohydride solution. The arrays are submerged in water for 2 min at 95°C, transferred into 0.2% SDS for 1 min, rinsed twice with water, air dried and stored in the dark at 25°C.

Cell or tissue mRNA is isolated or commercially obtained and probes are prepared by a single round of reverse transcription. Probes are hybridized to 1 cm² microarrays under a 14 x 14 mm glass coverslip for 6-12 hours at 60°C. Arrays are washed for 5 min at 25°C in low stringency wash buffer (1 x SSC/0.2% SDS), then for 10 min at room temperature in high stringency wash buffer (0.1 x SSC/0.2% SDS). Arrays are scanned in 0.1 x SSC using a fluorescence laser scanning device fitted with a custom filter set. Accurate differential expression measurements are obtained by taking the average of the ratios of two independent hybridizations.

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Quantitative analysis of the expression of genes may also be performed with full length cDNAs, extended cDNAs, 5' ESTs, or fragments thereof in complementary DNA arrays as described by Pietu et al.. (Genome Research 6:492-503, 1996). The full length cDNAs, extended cDNAs, 5' ESTs or fragments thereof are PCR amplified and spotted on membranes. Then, mRNAs originating from various tissues or cells are labeled with radioactive nucleotides. After hybridization and washing in controlled conditions, the hybridized mRNAs are detected by phospho-imaging or autoradiography. Duplicate experiments are performed and a quantitative analysis of differentially expressed mRNAs is then performed.

Alternatively, expression analysis of the 5' ESTs or extended cDNAs can be done through high density nucleotide arrays as described by Lockhart et al. (Nature Biotechnology 14: 1675-1680, 1996) and Sosnowsky et al. (Proc. Natl. Acad. Sci. 94:1119-1123, 1997). Oligonucleotides of 15-50 nucleotides corresponding to sequences of the 5' ESTs or extended cDNAs are synthesized directly on the chip (Lockhart et al., supra) or synthesized and then addressed to the chip (Sosnowsky et al., supra). Preferably, the oligonucleotides are about 20 nucleotides in length.

cDNA probes labeled with an appropriate compound, such as biotin, digoxigenin or fluorescent dye, are synthesized from the appropriate mRNA population and then randomly fragmented to an average size of 50 to 100 nucleotides. The said probes are then hybridized to the chip. After washing as described in Lockhart *et al*, *supra* and application of different electric fields (Sonowsky et *al*, *supra*.), the dyes or labeling compounds are detected and quantified. Duplicate hybridizations are performed. Comparative analysis of the intensity of the signal originating from cDNA probes on the same target oligonucleotide in different cDNA samples indicates a differential expression of the mRNA corresponding to the 5' EST or extended cDNA from which the oligonucleotide sequence has been designed.

III. Use of 5' ESTs to Clone Extended cDNAs and to Clone the Corresponding Genomic DNAs

Once 5' ESTs which include the 5' end of the corresponding mRNAs have been selected using the procedures described above, they can be utilized to isolate extended

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cDNAs which contain sequences adjacent to the 5' ESTs. The extended cDNAs may include the entire coding sequence of the protein encoded by the corresponding mRNA, including the authentic translation start site, the signal sequence, and the sequence encoding the mature protein remaining after cleavage of the signal peptide. Such extended cDNAs are referred to herein as "full length cDNAs." Alternatively, the extended cDNAs may include only the sequence encoding the mature protein remaining after cleavage of the signal peptide, or only the sequence encoding the signal peptide.

Example 27 below describes a general method for obtaining extended cDNAs using 5' ESTs. Example 28 below provides experimental results, using the method explained in example 27, describing several extended cDNAs including the entire coding sequence and authentic 5' end of the corresponding mRNA for several secreted proteins.

The methods of Examples 27, 28, and 29 can also be used to obtain extended cDNAs which encode less than the entire coding sequence of the secreted proteins encoded by the genes corresponding to the 5' ESTs. In some embodiments, the extended cDNAs isolated using these methods encode at least 10 amino acids of one of the proteins encoded by the sequences of SEQ ID NOs: 38-315. In further embodiments, the extended cDNAs encode at least 20 amino acids of the proteins encoded by the sequences of SEQ ID NOs: 38-315. In further embodiments, the extended cDNAs encode at least 30 amino amino acids of the sequences of SEQ ID NOs: 38-315. In a preferred embodiment, the extended cDNAs encode a full length protein sequence, which includes the protein coding sequences of SEQ ID NOs: 38-315.

EXAMPLE 27

General Method for Using 5' ESTs to Clone and Sequence cDNAs which Include the Entire Coding Region and the Authentic 5' End of the Corresponding mRNA

The following general method has been used to quickly and efficiently isolate extended cDNAs having the authentic 5' ends of their corresponding mRNAs as well as the full protein coding sequence and including sequence adjacent to the sequences of the 5' ESTs used to obtain them. This method may be applied to obtain extended cDNAs for any 5' EST in the NetGeneTM database, including those 5' ESTs encoding polypeptides belonging to secreted proteins. The method is summarized in figure 3.

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1. Obtention of Extended cDNAs

a) First strand synthesis

The method takes advantage of the known 5' sequence of the mRNA. A reverse transcription reaction is conducted on purified mRNA with a poly 14dT primer containing a 49 nucleotide sequence at its 5' end allowing the addition of a known sequence at the end of the cDNA which corresponds to the 3' end of the mRNA. For example, the primer may have the following sequence: 5'-ATC GTT GAG ACT CGT ACC AGC AGA GTC ACG AGA GAG ACT ACA CGG TAC TGG TTT TTT TTT TTT TTVN -3' (SEQ ID NO:14). Those skilled in the art will appreciate that other sequences may also be added to the poly dT sequence and used to prime the first strand synthesis. Using this primer and a reverse transcriptase such as the Superscript II (Gibco BRL) or Rnase H Minus M-MLV (Promega) enzyme, a reverse transcript anchored at the 3' polyA site of the RNAs is generated.

After removal of the mRNA hybridized to the first cDNA strand by alkaline hydrolysis, the products of the alkaline hydrolysis and the residual poly dT primer are eliminated with an exclusion column such as an AcA34 (Biosepra) matrix as explained in Example 11.

b) Second strand synthesis

A pair of nested primers on each end is designed based on the known 5' sequence from the 5' EST and the known 3' end added by the poly dT primer used in the first strand synthesis. Softwares used to design primers are either based on GC content and melting temperatures of oligonucleotides, such as OSP (Illier and Green, *PCR Meth. Appl.* 1:124-128, 1991), or based on the octamer frequency disparity method (Griffais *et al.*, *Nucleic Acids Res.* 19: 3887-3891, 1991) such as PC-Rare (http://bioinformatics.weizmann.ac.il/software/PC-Rare/doc/manuel.html).

Preferably, the nested primers at the 5' end are separated from one another by four to nine bases. The 5' primer sequences may be selected to have melting temperatures and specificities suitable for use in PCR.

Preferably, the nested primers at the 3' end are separated from one another by four to nine bases. For example, the nested 3' primers may have the following sequences: (5'- CCA GCA GAG TCA CGA GAG AGA CTA CAC GG -3'(SEQ ID NO:15), and 5'- CAC GAG AGA GAC TAC ACG GTA CTG G -3' (SEQ ID NO:16). These primers were selected

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because they have melting temperatures and specificities compatible with their use in PCR. However, those skilled in the art will appreciate that other sequences may also be used as primers.

The first PCR run of 25 cycles is performed using the Advantage Tth Polymerase Mix (Clontech) and the outer primer from each of the nested pairs. A second 20 cycle PCR using the same enzyme and the inner primer from each of the nested pairs is then performed on 1/2500 of the first PCR product. Thereafter, the primers and nucleotides are removed.

10 2. Sequencing of Full Length Extended cDNAs or Fragments Thereof

Due to the lack of position constraints on the design of 5' nested primers compatible for PCR use using the OSP software, amplicons of two types are obtained. Preferably, the second 5' primer is located upstream of the translation initiation codon thus yielding a nested PCR product containing the whole coding sequence. Such a full length extended cDNA undergoes a direct cloning procedure as described in section a. However, in some cases, the second 5' primer is located downstream of the translation initiation codon, thereby yielding a PCR product containing only part of the ORF. Such incomplete PCR products are submitted to a modified procedure described in section b. a) Nested PCR products containing complete ORFs

When the resulting nested PCR product contains the complete coding sequence, as predicted from the 5'EST sequence, it is cloned in an appropriate vector such as pED6dpc2, as described in section 3.

b) Nested PCR products containing incomplete ORFs

When the amplicon does not contain the complete coding sequence, intermediate steps are necessary to obtain both the complete coding sequence and a PCR product containing the full coding sequence. The complete coding sequence can be assembled from several partial sequences determined directly from different PCR products as described in the following section.

Once the full coding sequence has been completely determined, new primers compatible for PCR use are designed to obtain amplicons containing the whole coding region. However, in such cases, 3' primers compatible for PCR use are located inside the

3' UTR of the corresponding mRNA, thus yielding amplicons which lack part of this region, *i.e.* the polyA tract and sometimes the polyadenylation signal, as illustrated in figure 3. Such full length extended cDNAs are then cloned into an appropriate vector as described in section 3.

5 c) Sequencing extended cDNAs

Sequencing of extended cDNAs is performed using a Die Terminator approach with the AmpliTaq DNA polymerase FS kit available from Perkin Elmer.

In order to sequence PCR fragments, primer walking is performed using software such as OSP to choose primers and automated computer software such as ASMG (Sutton et al., Genome Science Technol. 1: 9-19, 1995) to construct contigs of walking sequences including the initial 5' tag using minimum overlaps of 32 nucleotides. Preferably, primer walking is performed until the sequences of full length cDNAs are obtained.

Completion of the sequencing of a given extended cDNA fragment is assessed as follows. Since sequences located after a polyA tract are difficult to determine precisely in the case of uncloned products, sequencing and primer walking processes for PCR products are interrupted when a polyA tract is identified in extended cDNAs obtained as described in case b. The sequence length is compared to the size of the nested PCR product obtained as described above. Due to the limited accuracy of the determination of the PCR product size by gel electrophoresis, a sequence is considered complete if the size of the obtained sequence is at least 70 % the size of the first nested PCR product. If the length of the sequence determined from the computer analysis is not at least 70% of the length of the nested PCR product, these PCR products are cloned and the sequence of the insertion is determined. When Northern blot data are available, the size of the mRNA detected for a given PCR product is used to finally assess that the sequence is complete. Sequences which do not fulfill the above criteria are discarded and will undergo a new isolation procedure.

Sequence data of all extended cDNAs are then transferred to a proprietary database, where quality controls and validation steps are carried out as described in example 15.

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3. Cloning of Full Length Extended cDNAs

The PCR product containing the full coding sequence is then cloned in an appropriate vector. For example, the extended cDNAs can be cloned into the expression vector pED6dpc2 (DiscoverEase, Genetics Institute, Cambridge, MA) as follows. pED6dpc2 vector DNA is prepared with blunt ends by performing an EcoRI digestion followed by a fill in reaction. The blunt ended vector is dephosphorylated. After removal of PCR primers and ethanol precipitation, the PCR product containing the full coding sequence or the extended cDNA obtained as described above is phosphorylated with a kinase subsequently removed by phenol-Sevag extraction and precipitation. The double stranded extended cDNA is then ligated to the vector and the resulting expression plasmid introduced into appropriate host cells.

Since the PCR products obtained as described above are blunt ended molecules that can be cloned in either direction, the orientation of several clones for each PCR product is determined. Then, 4 to 10 clones are ordered in microtiter plates and subjected to a PCR reaction using a first primer located in the vector close to the cloning site and a second primer located in the portion of the extended cDNA corresponding to the 3' end of the mRNA. This second primer may be the antisense primer used in anchored PCR in the case of direct cloning (case a) or the antisense primer located inside the 3'UTR in the case of indirect cloning (case b). Clones in which the start codon of the extended cDNA is operably linked to the promoter in the vector so as to permit expression of the protein encoded by the extended cDNA are conserved and sequenced. In addition to the ends of cDNA inserts, approximately 50 bp of vector DNA on each side of the cDNA insert are also sequenced.

The cloned PCR products are then entirely sequenced according to the aforementioned procedure. In this case, contigation of long fragments is then performed on walking sequences that have already contigated for uncloned PCR products during primer walking. Sequencing of cloned amplicons is complete when the resulting contigs include the whole coding region as well as overlapping sequences with vector DNA on both ends.

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4. Computer analysis of Full Length Extended cDNA

Sequences of all full length extended cDNAs are then submitted to further analysis as described below. Before searching the extended full length cDNAs for sequences of interest, extended cDNAs which are not of interest (vector RNAs, transfer RNAs, ribosomal RNAs, mitochondrial RNAs, prokaryotic RNAs and fungal RNAs) are discarded using methods essentially similar to those described for 5'ESTs in Example 18.

a) Identification of structural features

Structural features, e.g. polyA tail and polyadenylation signal, of the sequences of full length extended cDNAs are subsequently determined as follows.

A polyA tail is defined as a homopolymeric stretch of at least 11 A with at most one alternative base within it. The polyA tail search is restricted to the last 100 nt of the sequence and limited to stretches of 11 consecutive A's because sequencing reactions are often not readable after such a polyA stretch. Stretches having more than 90% homology over 8 nucleotides are identified as polyA tails using BLAST2N.

To search for a polyadenylation signal, the polyA tail is clipped from the full-length sequence. The 50 bp preceding the polyA tail are first searched for the canonic polyadenylation AAUAAA signal and, if the canonic signal is not detected, for the alternative AUUAAA signal (Sheets et al., Nuc. Acids Res. 18: 5799-5805, 1990). If neither of these consensus polyadenylation signals is found, the canonic motif is searched again allowing one mismatch to account for possible sequencing errors. More than 85 % of identified polyadenylation signals of either type actually ends 10 to 30 bp from the polyA tail. Alternative AUUAAA signals represents approximately 15 % of the total number of identified polyadenylation signals.

b) Identification of functional features

Functional features, e.g. ORFs and signal sequences, of the sequences of full length extended cDNAs were subsequently determined as follows.

The 3 upper strand frames of extended cDNAs are searched for ORFs defined as the maximum length fragments beginning with a translation intiation codon and ending with a stop codon. ORFs encoding at least 20 amino acids are preferred.

Each found ORF is then scanned for the presence of a signal peptide in the first 50 amino-acids or, where appropriate, within shorter regions down to 20 amino acids or

less in the ORF, using the matrix method of von Heijne (Nuc. Acids Res. 14: 4683-4690, 1986), the disclosure of which is incorporated herein by reference as described in Example 22.

c) Homology to either nucleotidic or proteic sequences

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Categorization of full-length sequences may be achieved using procedures essentially similar to those described for 5'ESTs in Example 24.

Extended cDNAs prepared as described above may be subsequently engineered to obtain nucleic acids which include desired portions of the extended cDNA using conventional techniques such as subcloning, PCR, or *in vitro* oligonucleotide synthesis. For example, nucleic acids which include only the full coding sequences (*i.e.* the sequences encoding the signal peptide and the mature protein remaining after the signal peptide is cleaved off) may be obtained using techniques known to those skilled in the art. Alternatively, conventional techniques may be applied to obtain nucleic acids which contain only the coding sequences for the mature protein remaining after the signal peptide is cleaved off or nucleic acids which contain only the coding sequences for the signal peptides.

Similarly, nucleic acids containing any other desired portion of the coding sequences for the secreted protein may be obtained. For example, the nucleic acid may contain at least 10 consecutive bases of an extended cDNA such as one of the extended cDNAs described below. In another embodiment, the nucleic acid may contain at least 15 consecutive bases of an extended cDNA such as one of the extended cDNAs described below. Alternatively, the nucleic acid may contain at least 20 consecutive bases of an extended cDNA such as one of the extended cDNAs described below. In another embodiment, the nucleic acid may contain at least 25 consecutive bases of an extended cDNAs uch as one of the extended cDNAs described below. In yet another embodiment, the nucleic acid may contain at least 40 consecutive bases of an extended cDNA such as one of the extended cDNAs described below.

Once an extended cDNA has been obtained, it can be sequenced to determine the amino acid sequence it encodes. Once the encoded amino acid sequence has been determined, one can create and identify any of the many conceivable cDNAs that will encode that protein by simply using the degeneracy of the genetic code. For example, allelic variants

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or other homologous nucleic acids can be identified as described below. Alternatively, nucleic acids encoding the desired amino acid sequence can be synthesized *in vitro*.

In a preferred embodiment, the coding sequence may be selected using the known codon or codon pair preferences for the host organism in which the cDNA is to be expressed.

The extended cDNAs derived from the 5' ESTS of the present invention were obtained as described in Example 28 below.

EXAMPLE 28

Characterization of cloned extended cDNAs obtained using 5' ESTs

The procedure described in Example 27 above was used to obtain the extended cDNAs derived from the 5' ESTs of the present invention in a variety of tissues. The following list provides a few examples of thus obtained extended cDNAs.

Using this approach, the full length cDNA of SEQ ID NO:17 (internal identification number 48-19-3-G1-FL1) was obtained. This cDNA falls into the "EST-ext" category described above and encodes the signal peptide MKKVLLLITAILAVAVG (SEQ ID NO: 18) having a von Heijne score of 8.2.

The full length cDNA of SEQ ID NO:19 (internal identification number 58-34-2-E7-FL2) was also obtained using this procedure. This cDNA falls into the "EST-ext" category described above and encodes the signal peptide MWWFQQGLSFLPSALVIWTSA (SEQ ID NO:20) having a von Heijne score of 5.5.

Another full length cDNA obtained using the procedure described above has the sequence of SEQ ID NO:21 (internal identification number 51-27-1-E8-FL1). This cDNA, falls into the "EST-ext" category described above and encodes the signal peptide MVLTTLPSANSANSPVNMPTTGPNSLSYASSALSPCLT (SEQ ID NO:22) having a von Heijne score of 5.9.

The above procedure was also used to obtain a full length cDNA having the sequence of SEQ ID NO:23 (internal identification number 76-4-1-G5-FL1). This cDNA falls into the "EST-ext" category described above and encodes the signal peptide ILSTVTALTFAXA (SEQ ID NO:24) having a von Heijne score of 5.5.

The full length cDNA of SEQ ID NO:25 (internal identification number 51-3-3-B10-FL3) was also obtained using this procedure. This cDNA falls into the "new" category

described above and encodes a signal peptide LVLTLCTLPLAVA (SEQ ID NO:26) having a von Heijne score of 10.1.

The full length cDNA of SEQ ID NO.27 (internal identification number 58-35-2-F10-FL2) was also obtained using this procedure. This cDNA falls into the "new" category described above and encodes a signal peptide LWLLFFLVTAIHA (SEQ ID NO:28) having a von Heijne score of 10.7.

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Bacterial clones containing plasmids containing the full length cDNAs described above are presently stored in the inventor's laboratories under the internal identification numbers provided above. The inserts may be recovered from the stored materials by growing an aliquot of the appropriate bacterial clone in the appropriate medium. The plasmid DNA can then be isolated using plasmid isolation procedures familiar to those skilled in the art such as alkaline lysis minipreps or large scale alkaline lysis plasmid isolation procedures. If desired the plasmid DNA may be further enriched by centrifugation on a cesium chloride gradient, size exclusion chromatography, or anion exchange chromatography. The plasmid DNA obtained using these procedures may then be manipulated using standard cloning techniques familiar to those skilled in the art. Alternatively, a PCR can be done with primers designed at both ends of the cDNA insertion. The PCR product which corresponds to the cDNA can then be manipulated using standard cloning techniques familiar to those skilled in the art.

The polypeptides encoded by the extended cDNAs may be screened for the presence of known structural or functional motifs or for the presence of signatures, small amino acid sequences which are well conserved amongst the members of a protein family. The conserved regions have been used to derive consensus patterns or matrices included in the PROSITE data bank, in particular in the file prosite.dat (Release 13.0 of November 1995, located at http://expasy.hcuge.ch/sprot/prosite.html. Prosite_convert and prosite_scan programs (http://ulrec3.unil.ch/ftpserveur/prosite_scan) may be used to find signatures on the extended cDNAs.

For each pattern obtained with the prosite_convert program from the prosite dat file, the accuracy of the detection on a new protein sequence may be assessed by evaluating the frequency of irrelevant hits on the population of human secreted proteins included in the data bank SWISSPROT. The ratio between the number of hits on shuffled proteins (with a window size of 20 amino acids) and the number of hits on native (unshuffled) proteins may be

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used as an index. Every pattern for which the ratio is greater than 20% (one hit on shuffled proteins for 5 hits on native proteins) may be skipped during the search with prosite_scan. The program used to shuffle protein sequences (db_shuffled) and the program used to determine the statistics for each pattern in the protein data banks (prosite_statistics) are available on the ftp site http://ulrec3.unil.ch/ftpserveur/prosite_scan.

In addition to PCR based methods for obtaining extended cDNAs, traditional hybridization based methods may also be employed. These methods may also be used to obtain the genomic DNAs which encode the mRNAs from which the 5' ESTs were derived, mRNAs corresponding to the extended cDNAs, or nucleic acids which are homologous to extended cDNAs or 5' ESTs. Example 29 below provides examples of such methods.

EXAMPLE 29

Methods for Obtaining cDNAs which include the Entire Coding Region and the Authentic 5'End of the Corresponding mRNA

A full length cDNA library can be made using the strategies described in Examples 13, 14, 15, and 16 above by replacing the random nonamer used in Example 14 with an oligo-dT primer. For instance, the oligonucleotide of SEQ ID NO:14 may be used.

Alternatively, a cDNA library or genomic DNA library may be obtained from a commercial source or made using techniques familiar to those skilled in the art. Such cDNA or genomic DNA librairies may be used to isolate extended cDNAs obtained from 5' EST or nucleic acids homologous to extended cDNAs or 5' EST as follows. The cDNA library or genomic DNA library is hybridized to a detectable probe comprising at least 10 consecutive nucleotides from the 5' EST or extended cDNA using conventional techniques. Preferably, the probe comprises at least 12, 15, or 17 consecutive nucleotides from the 5' EST or extended cDNA. More preferably, the probe comprises at least 20 to 30 consecutive nucleotides from the 5' EST or extended cDNA. In some embodiments, the probe comprises more than 30 nucleotides from the 5' EST or extended cDNA.

Techniques for identifying cDNA clones in a cDNA library which hybridize to a given probe sequence are disclosed in Sambrook et al., Molecular Cloning: A Laboratory Manual

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2d Ed., Cold Spring Harbor Laboratory Press, 1989, the disclosure of which is incorporated herein by reference. The same techniques may be used to isolate genomic DNAs.

Briefly, cDNA or genomic DNA clones which hybridize to the detectable probe are identified and isolated for further manipulation as follows. A probe comprising at least 10 consecutive nucleotides from the 5' EST or extended cDNA is labeled with a detectable label such as a radioisotope or a fluorescent molecule. Preferably, the probe comprises at least 12, 15, or 17 consecutive nucleotides from the 5' EST or extended cDNA. More preferably, the probe comprises 20 to 30 consecutive nucleotides from the 5' EST or extended cDNA. In some embodiments, the probe comprises more than 30 nucleotides from the 5' EST or extended cDNA.

Techniques for labeling the probe are well known and include phosphorylation with polynucleotide kinase, nick translation, *in vitro* transcription, and non radioactive techniques. The cDNAs or genomic DNAs in the library are transferred to a nitrocellulose or nylon filter and denatured. After blocking of non specific sites, the filter is incubated with the labeled probe for an amount of time sufficient to allow binding of the probe to cDNAs or genomic DNAs containing a sequence capable of hybridizing thereto.

By varying the stringency of the hybridization conditions used to identify extended cDNAs or genomic DNAs which hybridize to the detectable probe, extended cDNAS having different levels of homology to the probe can be identified and isolated as described below.

1. Identification of Extended cDNA or Genomic cDNA Sequences Having a High Degree of Homology to the Labeled Probe

To identify extended cDNAs or genomic DNAs having a high degree of homology to the probe sequence, the melting temperature of the probe may be calculated using the following formulas:

For probes between 14 and 70 nucleotides in length the melting temperature (Tm) is calculated using the formula: Tm=81.5+16.6(log [Na+])+0.41(fraction G+C)-(600/N) where N is the length of the probe.

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If the hybridization is carried out in a solution containing formamide, the melting temperature may be calculated using the equation Tm=81.5+16.6(log [Na+])+0.41(fraction G+C)-(0.63% formamide)-(600/N) where N is the length of the probe.

Prehybridization may be carried out in 6X SSC, 5X Denhardt's reagent, 0.5% SDS, 100 µg denatured fragmented salmon sperm DNA or 6X SSC, 5X Denhardt's reagent, 0.5% SDS, 100 µg denatured fragmented salmon sperm DNA, 50% formamide. The formulas for SSC and Denhardt's solutions are listed in Sambrook *et al.*, *supra*.

Hybridization is conducted by adding the detectable probe to the prehybridization solutions listed above. Where the probe comprises double stranded DNA, it is denatured before addition to the hybridization solution. The filter is contacted with the hybridization solution for a sufficient period of time to allow the probe to hybridize to extended cDNAs or genomic DNAs containing sequences complementary thereto or homologous thereto. For probes over 200 nucleotides in length, the hybridization may be carried out at 15-25°C below the Tm. For shorter probes, such as oligonucleotide probes, the hybridization may be conducted at 15-25°C below the Tm. Preferably, for hybridizations in 6X SSC, the hybridization is conducted at approximately 68°C. Preferably, for hybridizations in 50% formamide containing solutions, the hybridization is conducted at approximately 42°C.

All of the foregoing hybridizations would be considered to be under "stringent" conditions.

Following hybridization, the filter is washed in 2X SSC, 0.1% SDS at room temperature for 15 minutes. The filter is then washed with 0.1X SSC, 0.5% SDS at room temperature for 30 minutes to 1 hour. Thereafter, the solution is washed at the hybridization temperature in 0.1X SSC, 0.5% SDS. A final wash is conducted in 0.1X SSC at room temperature.

Extended cDNAs, nucleic acids homologous to extended cDNAs or 5' ESTs, or genomic DNAs which have hybridized to the probe are identified by autoradiography or other conventional techniques.

2. Obtention of Extended cDNA or Genomic cDNA Sequences Having Lower Degrees of Homology to the Labeled Probe

The above procedure may be modified to identify extended cDNAs, nucleic acids homologous to extended cDNAs, or genomic DNAs having decreasing levels of homology to the probe sequence. For example, to obtain extended cDNAs, nucleic acids homologous to extended cDNAs, or genomic DNAs of decreasing homology to the detectable probe, less stringent conditions may be used. For example, the hybridization temperature may be decreased in increments of 5°C from 68°C to 42°C in a hybridization buffer having a sodium concentration of approximately 1M. Following hybridization, the filter may be washed with 2X SSC, 0.5% SDS at the temperature of hybridization. These conditions are considered to be "moderate" conditions above 50°C and "low" conditions below 50°C.

Alternatively, the hybridization may be carried out in buffers, such as 6X SSC, containing formamide at a temperature of 42°C. In this case, the concentration of formamide in the hybridization buffer may be reduced in 5% increments from 50% to 0% to identify clones having decreasing levels of homology to the probe. Following hybridization, the filter may be washed with 6X SSC, 0.5% SDS at 50°C. These conditions are considered to be "moderate" conditions above 25% formamide and "low" conditions below 25% formamide.

Extended cDNAs, nucleic acids homologous to extended cDNAs, or genomic DNAs which have hybridized to the probe are identified by autoradiography.

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3. Determination of the Degree of Homology Between the Obtained Extended cDNAs and the Labeled Probe

If it is desired to obtain nucleic acids homologous to extended cDNAs, such as allelic variants thereof or nucleic acids encoding proteins related to the proteins encoded by the extended cDNAs, the level of homology between the hybridized nucleic acid and the extended cDNA or 5' EST used as the probe may be further determined using BLAST2N; parameters may be adapted depending on the sequence length and degree of homology studied. To determine the level of homology between the hybridized nucleic acid and the extended cDNA or 5'EST from which the probe was derived, the nucleotide sequences of the hybridized nucleic acid and the extended cDNA or 5'EST from which the probe was derived are compared. For example, using the above methods, nucleic acids having at least 95%

nucleic acid homology to the extended cDNA or 5'EST from which the probe was derived may be obtained and identified. Similarly, by using progressively less stringent hybridization conditions one can obtain and identify nucleic acids having at least 90%, at least 85%, at least 80% or at least 75% homology to the extended cDNA or 5'EST from which the probe was derived.

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To determine whether a clone encodes a protein having a given amount of homology to the protein encoded by the extended cDNA or 5' EST, the amino acid sequence encoded by the extended cDNA or 5' EST is compared to the amino acid sequence encoded by the hybridizing nucleic acid. Homology is determined to exist when an amino acid sequence in the extended cDNA or 5' EST is closely related to an amino acid sequence in the hybridizing nucleic acid. A sequence is closely related when it is identical to that of the extended cDNA or 5' EST or when it contains one or more amino acid substitutions therein in which amino acids having similar characteristics have been substituted for one another. Using the above methods and algorithms such as FASTA with parameters depending on the sequence length and degree of homology studied, one can obtain nucleic acids encoding proteins having at least 95%, at least 90%, at least 85%, at least 80% or at least 75% homology to the proteins encoded by the extended cDNA or 5'EST from which the probe was derived.

In addition to the above described methods, other protocols are available to obtain extended cDNAs using 5' ESTs as outlined in the following paragraphs.

Extended cDNAs may be prepared by obtaining mRNA from the tissue, cell, or organism of interest using mRNA preparation procedures utilizing polyA selection procedures or other techniques known to those skilled in the art. A first primer capable of hybridizing to the polyA tail of the mRNA is hybridized to the mRNA and a reverse transcription reaction is performed to generate a first cDNA strand.

The first cDNA strand is hybridized to a second primer containing at least 10 consecutive nucleotides of the sequences of SEQ ID NOs 38-315. Preferably, the primer comprises at least 12, 15, or 17 consecutive nucleotides from the sequences of SEQ ID NOs 38-315. More preferably, the primer comprises 20 to 30 consecutive nucleotides from the sequences of SEQ ID NOs 38-315. In some embodiments, the primer comprises more than 30 nucleotides from the sequences of SEQ ID NOs 38-315. If it is desired to obtain extended

cDNAs containing the full protein coding sequence, including the authentic translation initiation site, the second primer used contains sequences located upstream of the translation initiation site. The second primer is extended to generate a second cDNA strand complementary to the first cDNA strand. Alternatively, RT-PCR may be performed as described above using primers from both ends of the cDNA to be obtained.

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Extended cDNAs containing 5' fragments of the mRNA may be prepared by hybridizing an mRNA comprising the sequence of the 5'EST for which an extended cDNA is desired with a primer comprising at least 10 consecutive nucleotides of the sequences complementary to the 5'EST and reverse transcribing the hybridized primer to make a first cDNA strand from the mRNAs. Preferably, the primer comprises at least 12, 15, or 17 consecutive nucleotides from the 5'EST. More preferably, the primer comprises 20 to 30 consecutive nucleotides from the 5'EST.

Thereafter, a second cDNA strand complementary to the first cDNA strand is synthesized. The second cDNA strand may be made by hybridizing a primer complementary to sequences in the first cDNA strand to the first cDNA strand and extending the primer to generate the second cDNA strand.

The double stranded extended cDNAs made using the methods described above are isolated and cloned. The extended cDNAs may be cloned into vectors such as plasmids or viral vectors capable of replicating in an appropriate host cell. For example, the host cell may be a bacterial, mammalian, avian, or insect cell.

Techniques for isolating mRNA, reverse transcribing a primer hybridized to mRNA to generate a first cDNA strand, extending a primer to make a second cDNA strand complementary to the first cDNA strand, isolating the double stranded cDNA and cloning the double stranded cDNA are well known to those skilled in the art and are described in Current Protocols in Molecular Biology, John Wiley and Sons, Inc. 1997 and Sambrook et al., Molecular Cloning: A Laboratory Manual, Second Edition, Cold Spring Harbor Laboratory Press, 1989, the entire disclosures of which are incorporated herein by reference.

Alternatively, procedures such as the one described in Example 29 may be used for obtaining full length cDNAs or extended cDNAs. In this approach, full length or extended cDNAs are prepared from mRNA and cloned into double stranded phagemids as follows. The cDNA library in the double stranded phagemids is then rendered single stranded by

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treatment with an endonuclease, such as the Gene II product of the phage F1, and an exonuclease (Chang et al., Gene 127:95-8, 1993). A biotinylated oligonucleotide comprising the sequence of a 5' EST, or a fragment containing at least 10 nucleotides thereof, is hybridized to the single stranded phagemids. Preferably, the fragment comprises at least 12, 15, or 17 consecutive nucleotides from the 5' EST. More preferably, the fragment comprises 20-30 consecutive nucleotides from the 5' EST. In some procedures, the fragment may comprise more than 30 consecutive nucleotides from the 5' EST.

Hybrids between the biotinylated oligonucleotide and phagemids having inserts containing the 5' EST sequence are isolated by incubating the hybrids with streptavidin coated paramagnetic beads and retrieving the beads with a magnet (Fry et al., Biotechniques, 13: 124-131, 1992). Therafter, the resulting phagemids containing the 5' EST sequence are released from the beads and converted into double stranded DNA using a primer specific for the 5' EST sequence. Alternatively, protocoles such as the Gene Trapper kit (Gibco BRL) may be used. The resulting double stranded DNA is transformed into bacteria. Extended cDNAs containing the 5' EST sequence are identified by colony PCR or colony hybridization.

Using any of the above described methods in section III, a plurality of extended cDNAs containing full length protein coding sequences or sequences encoding only the mature protein remaining after the signal peptide is cleaved off may be provided as cDNA libraries for subsequent evaluation of the encoded proteins or use in diagnostic assays as described below.

IV. Expression of Proteins Encoded by Extended cDNAs Isolated Using 5' ESTs

Extended cDNAs containing the full protein coding sequences of their corresponding mRNAs or portions thereof, such as cDNAs encoding the mature protein, may be used to express the encoded secreted proteins or portions thereof as described in Example 30 below. If desired, the extended cDNAs may contain the sequences encoding the signal peptide to facilitate secretion of the expressed protein. It will be appreciated that a plurality of extended cDNAs containing the full protein coding sequences or portions thereof may be simultaneously cloned into expression vectors to create an expression library for analysis of the encoded proteins as described below.

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EXAMPLE 30

Expression of the Proteins Encoded by the Genes Corresponding to 5'ESTS or Portions Thereof

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To express the proteins encoded by the genes corresponding to 5' ESTs (or portions thereof), full length cDNAs containing the entire protein coding region or extended cDNAs containing sequences adjacent to the 5' ESTs (or portions thereof) are obtained as described in Examples 27-29 and cloned into a suitable expression vector. If desired, the nucleic acids may contain the sequences encoding the signal peptide to facilitate secretion of the expressed protein. The nucleic acids inserted into the expression vectors may also contain sequences upstream of the sequences encoding the signal peptide, such as sequences which regulate expression levels or sequences which confer tissue specific expression.

The nucleic acid encoding the protein or polypeptide to be expressed is operably linked to a promoter in an expression vector using conventional cloning technology. The expression vector may be any of the mammalian, yeast, insect or bacterial expression systems known in the art. Commercially available vectors and expression systems are available from a variety of suppliers including Genetics Institute (Cambridge, MA), Stratagene (La Jolla, California), Promega (Madison, Wisconsin), and Invitrogen (San Diego, California). If desired, to enhance expression and facilitate proper protein folding, the codon context and codon pairing of the sequence may be optimized for the particular expression organism in which the expression vector is introduced, as explained by Hatfield, *et al.*, U.S. Patent No. 5,082,767, incorporated herein by this reference.

The cDNA cloned into the expression vector may encode the entire protein (i.e. the signal peptide and the mature protein), the mature protein (i.e. the protein created by cleaving the signal peptide off), only the signal peptide or any other portion thereof.

The following is provided as one exemplary method to express the proteins encoded by the extended cDNAs corresponding to the 5' ESTs or the nucleic acids described above. First, the methionine initiation codon for the gene and the polyA signal of the gene are identified. If the nucleic acid encoding the polypeptide to be expressed lacks a methionine to serve as the initiation site, an initiating methionine can be introduced next to the first codon of the nucleic acid using conventional techniques. Similarly, if the extended cDNA lacks a polyA signal, this sequence can be added to the construct by, for example, splicing out the

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polyA signal from pSG5 (Stratagene) using BglII and SalI restriction endonuclease enzymes and incorporating it into the mammalian expression vector pXT1 (Stratagene). pXT1 contains the LTRs and a portion of the gag gene from Moloney Murine Leukemia Virus. The position of the LTRs in the construct allow efficient stable transfection. The vector includes the Herpes Simplex thymidine kinase promoter and the selectable neomycin gene. The extended cDNA or portion thereof encoding the polypeptide to be expressed is obtained by PCR from the bacterial vector using oligonucleotide primers complementary to the extended cDNA or portion thereof and containing restriction endonuclease sequences for Pst I incorporated into the 5'primer and BglII at the 5' end of the corresponding cDNA 3' primer, taking care to ensure that the extended cDNA is positioned with the poly A signal. The purified fragment obtained from the resulting PCR reaction is digested with PstI, blunt ended with an exonuclease, digested with Bgl II, purified and ligated to pXT1 containing a poly A signal and prepared for this ligation (blunt/BglII).

The ligated product is transfected into mouse NIH 3T3 cells using Lipofectin (Life Technologies, Inc., Grand Island, New York) under conditions outlined in the product specification. Positive transfectants are selected after growing the transfected cells in 600 µg/ml G418 (Sigma, St. Louis, Missouri). Preferably the expressed protein is released into the culture medium, thereby facilitating purification.

Alternatively, the extended cDNAs may be cloned into pED6dpc2 as described above. The resulting pED6dpc2 constructs may be transfected into a suitable host cell, such as COS 1 cells. Methotrexate resistant cells are selected and expanded. Preferably, the protein expressed from the extended cDNA is released into the culture medium thereby facilitating purification.

Proteins in the culture medium are separated by gel electrophoresis. If desired, the proteins may be ammonium sulfate precipitated or separated based on size or charge prior to electrophoresis.

As a control, the expression vector lacking a cDNA insert is introduced into host cells or organisms and the proteins in the medium are harvested. The secreted proteins present in the medium are detected using techniques familiar to those skilled in the art such as Coomassie blue or silver staining or using antibodies against the protein encoded by the extended cDNA

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Antibodies capable of specifically recognizing the protein of interest may be generated using synthetic 15-mer peptides having a sequence encoded by the appropriate 5' EST, extended cDNA, or portion thereof. The synthetic peptides are injected into mice to generate antibody to the polypeptide encoded by the 5' EST, extended cDNA, or portion thereof.

Secreted proteins from the host cells or organisms containing an expression vector which contains the extended cDNA derived from a 5' EST or a portion thereof are compared to those from the control cells or organism. The presence of a band in the medium from the cells containing the expression vector which is absent in the medium from the control cells indicates that the extended cDNA encodes a secreted protein. Generally, the band corresponding to the protein encoded by the extended cDNA will have a mobility near that expected based on the number of amino acids in the open reading frame of the extended cDNA. However, the band may have a mobility different than that expected as a result of modifications such as glycosylation, ubiquitination, or enzymatic cleavage.

Alternatively, if the protein expressed from the above expression vectors does not contain sequences directing its secretion, the proteins expressed from host cells containing an expression vector with an insert encoding a secreted protein or portion thereof can be compared to the proteins expressed in control host cells containing the expression vector without an insert. The presence of a band in samples from cells containing the expression vector with an insert which is absent in samples from cells containing the expression vector without an insert indicates that the desired protein or portion thereof is being expressed. Generally, the band will have the mobility expected for the secreted protein or portion thereof. However, the band may have a mobility different than that expected as a result of modifications such as glycosylation, ubiquitination, or enzymatic cleavage.

The protein encoded by the extended cDNA may be purified using standard immunochromatography techniques. In such procedures, a solution containing the secreted protein, such as the culture medium or a cell extract, is applied to a column having antibodies against the secreted protein attached to the chromatography matrix. The secreted protein is allowed to bind the immunochromatography column. Thereafter, the column is washed to remove non-specifically bound proteins. The specifically bound secreted protein is then released from the column and recovered using standard techniques.

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If antibody production is not possible, the extended cDNA sequence or portion thereof may be incorporated into expression vectors designed for use in purification schemes employing chimeric polypeptides. In such strategies, the coding sequence of the extended cDNA or portion thereof is inserted in frame with the gene encoding the other half of the chimera. The other half of the chimera may be β -globin or a nickel binding polypeptide. A chromatography matrix having antibody to β -globin or nickel attached thereto is then used to purify the chimeric protein. Protease cleavage sites may be engineered between the β -globin gene or the nickel binding polypeptide and the extended cDNA or portion thereof. Thus, the two polypeptides of the chimera may be separated from one another by protease digestion.

One useful expression vector for generating β-globin chimerics is pSG5 (Stratagene), which encodes rabbit β-globin. Intron II of the rabbit β-globin gene facilitates splicing of the expressed transcript, and the polyadenylation signal incorporated into the construct increases the level of expression. These techniques as described are well known to those skilled in the art of molecular biology. Standard methods are published in methods texts such as Davis *et al.*, (*Basic Methods in Molecular Biology*, Davis, Dibner, and Battey, ed., Elsevier Press, NY, 1986) and many of the methods are available from Stratagene, Life Technologies, Inc., or Promega. Polypeptide may additionally be produced from the construct using *in vitro* translation systems such as the *In vitro* ExpressTM Translation Kit (Stratagene).

Following expression and purification of the secreted proteins encoded by the 5' ESTs, extended cDNAs, or fragments thereof, the purified proteins may be tested for the ability to bind to the surface of various cell types as described in Example 31 below. It will be appreciated that a plurality of proteins expressed from these cDNAs may be included in a panel of proteins to be simultaneously evaluated for the activities specifically described below, as well as other biological roles for which assays for determining activity are available.

EXAMPLE 31

Analysis of Secreted Proteins to Determine Whether they Bind to the Cell Surface

The proteins encoded by the 5' ESTs, extended cDNAs, or fragments thereof are cloned into expression vectors such as those described in Example 30. The proteins are purified by size, charge, immunochromatography or other techniques familiar to those skilled

in the art. Following purification, the proteins are labeled using techniques known to those skilled in the art. The labeled proteins are incubated with cells or cell lines derived from a variety of organs or tissues to allow the proteins to bind to any receptor present on the cell surface. Following the incubation, the cells are washed to remove non-specifically bound protein. The labeled proteins are detected by autoradiography. Alternatively, unlabeled proteins may be incubated with the cells and detected with antibodies having a detectable label, such as a fluorescent molecule, attached thereto.

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Specificity of cell surface binding may be analyzed by conducting a competition analysis in which various amounts of unlabeled protein are incubated along with the labeled protein. The amount of labeled protein bound to the cell surface decreases as the amount of competitive unlabeled protein increases. As a control, various amounts of an unlabeled protein unrelated to the labeled protein is included in some binding reactions. The amount of labeled protein bound to the cell surface does not decrease in binding reactions containing increasing amounts of unrelated unlabeled protein, indicating that the protein encoded by the cDNA binds specifically to the cell surface.

As discussed above, secreted proteins have been shown to have a number of important physiological effects and, consequently, represent a valuable therapeutic resource. The secreted proteins encoded by the extended cDNAs or portions thereof made according to Examples 27-29 may be evaluated to determine their physiological activities as described below.

EXAMPLE 32

Assaying the Proteins Expressed from Extended cDNAs or Portions Thereof for Cytokine, Cell Proliferation or Cell Differentiation Activity

As discussed above, secreted proteins may act as cytokines or may affect cellular proliferation or differentiation. Many protein factors discovered to date, including all known cytokines, have exhibited activity in one or more factor dependent cell proliferation assays, and hence the assays serve as a convenient confirmation of cytokine activity. The activity of a protein encoded by the extended cDNAs is evidenced by any one of a number of routine factor dependent cell proliferation assays for cell lines including, without limitation, 32D,

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DA2, DA1G, T10, B9, B9/11, BaF3, MC9/G, M⁺ (preB M⁺), 2E8, RB5, DA1, 123, T1165, HT2, CTLL2, TF-1, Mo7c and CMK. The proteins encoded by the above extended cDNAs or portions thereof may be evaluated for their ability to regulate T cell or thymocyte proliferation in assays such as those described above or in the following references, which are incorporated herein by reference: Current Protocols in Immunology, Ed. by Coligan et al., Greene Publishing Associates and Wiley-Interscience; Takai et al. J. Immunol. 137:3494-3500, 1986., Bertagnolli et al., J. Immunol. 145:1706-1712, 1990., Bertagnolli et al., Cell. Immunol. 133:327-341, 1991; Bertagnolli, et al., J. Immunol. 149:3778-3783, 1992; Bowman et al., J. Immunol. 152:1756-1761, 1994.

In addition, numerous assays for cytokine production and/or the proliferation of spleen cells, lymph node cells and thymocytes are known. These include the techniques disclosed in *Current Protocols in Immunology, supra* 1:3.12.1-3.12.14; and Schreiber In *Current Protocols in Immunology, supra* 1:6.8.1-6.8.8.

The proteins encoded by the cDNAs may also be assayed for the ability to regulate the proliferation and differentiation of hematopoietic or lymphopoietic cells. Many assays for such activity are familiar to those skilled in the art, including the assays in the following references, which are incorporated herein by reference: Bottomly et al., In Current Protocols in Immunology., supra. 1:6.3.1-6.3.12,; deVries et al., J. Exp. Med. 173:1205-1211, 1991; Moreau et al., Nature 36:690-692, 1988; Greenberger et al., Proc. Natl. Acad. Sci. U.S.A. 80:2931-2938, 1983; Nordan, R., In Current Protocols in Immunology., supra. 1:6.6.1-6.6.5; Smith et al., Proc. Natl. Acad. Sci. U.S.A. 83:1857-1861, 1986; Bennett et al., in Current Protocols in Immunology supra 1:6.15.1; Ciarletta et al., In Current Protocols in Immunology. supra 1:6.13.1.

The proteins encoded by the cDNAs may also be assayed for their ability to regulate T-cell responses to antigens. Many assays for such activity are familiar to those skilled in the art, including the assays described in the following references, which are incorporated herein by reference: Chapter 3 (*In Vitro* Assays for Mouse Lymphocyte Function), Chapter 6 (Cytokines and Their Cellular Receptors) and Chapter 7, (Immunologic Studies in Humans) in *Current Protocols in Immunology supra*; Weinberger et al., Proc. Natl. Acad. Sci. USA 77:6091-6095, 1980; Weinberger et al., Eur. J. Immun. 11:405-411, 1981; Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988.

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Those proteins which exhibit cytokine, cell proliferation, or cell differentiation activity may then be formulated as pharmaceuticals and used to treat clinical conditions in which induction of cell proliferation or differentiation is beneficial. Alternatively, as described in more detail below, genes encoding these proteins or nucleic acids regulating the expression of these proteins may be introduced into appropriate host cells to increase or decrease the expression of the proteins as desired.

EXAMPLE 33

Assaying the Proteins Expressed from Extended cDNAs or Portions

Thereof for Activity as Immune System Regulators

The proteins encoded by the cDNAs may also be evaluated for their effects as immune regulators. For example, the proteins may be evaluated for their activity to influence thymocyte or splenocyte cytotoxicity. Numerous assays for such activity are familiar to those skilled in the art including the assays described in the following references, which are incorporated herein by reference: Chapter 3 (In Vitro Assays for Mouse Lymphocyte Function 3.1-3.19) and Chapter 7 (Immunologic studies in Humans) in Current Protocols in Immunology, Coligan et al., Eds, Greene Publishing Associates and Wiley-Interscience; Herrmann et al., Proc. Natl. Acad. Sci. USA 78:2488-2492, 1981; Herrmann et al., J. Immunol. 128:1968-1974, 1982, Handa et al., J. Immunol. 135:1564-1572, 1985; Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Bowman et al., J. Virology 61:1992-1998; Bertagnolli et al., Cell. Immunol. 133:327-341, 1991; Brown et al., J. Immunol. 153:3079-3092, 1994.

The proteins encoded by the cDNAs may also be evaluated for their effects on T-cell dependent immunoglobulin responses and isotype switching. Numerous assays for such activity are familiar to those skilled in the art, including the assays disclosed in the following references, which are incorporated herein by reference: Maliszewski, *J. Immunol.* 144:3028-3033, 1990; Mond *et al.* in *Current Protocols in Immunology*, 1:3.8.1-3.8.16, *supra*.

The proteins encoded by the cDNAs may also be evaluated for their effect on immune effector cells, including their effect on Th1 cells and cytotoxic lymphocytes. Numerous assays for such activity are familiar to those skilled in the art, including the assays disclosed in the

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following references, which are incorporated herein by reference: Chapter 3 (In Vitro Assays for Mouse Lymphocyte Function 3.1-3.19) and Chapter 7 (Immunologic Studies in Humans) in Current Protocols in Immunology, supra; Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Bertagnolli et al., J. Immunol. 149:3778-3783, 1992.

The proteins encoded by the cDNAs may also be evaluated for their effect on dendritic cell mediated activation of naive T-cells. Numerous assays for such activity are familiar to those skilled in the art, including the assays disclosed in the following references, which are incorporated herein by reference: Guery et al., J. Immunol. 134:536-544, 1995; Inaba et al., J. Exp. Med. 173:549-559, 1991; Macatonia et al., J. Immunol. 154:5071-5079, 1995; Porgador et al.J. Exp. Med. 182:255-260, 1995; Nair et al., J. Virol. 67:4062-4069, 1993; Huang et al., Science 264:961-965, 1994; Macatonia et al.J. Exp. Med. 169:1255-1264, 1989; Bhardwaj et al., Journal of Clinical Investigation 94:797-807, 1994; and Inaba et al., J. Exp. Med. 172:631-640, 1990.

The proteins encoded by the cDNAs may also be evaluated for their influence on the lifetime of lymphocytes. Numerous assays for such activity are familiar to those skilled in the art, including the assays disclosed in the following references, which are incorporated herein by reference: Darzynkiewicz et al., Cytometry 13:795-808, 1992; Gorczyca et al., Leukemia 7:659-670, 1993; Gorczyca et al., Cancer Res. 53:1945-1951, 1993; Itoh et al., Cell 66:233-243, 1991; Zacharchuk, J. Immunol. 145:4037-4045, 1990; Zamai et al., Cytometry 14:891-897, 1993; Gorczyca et al., Int. J. Oncol. 1:639-648, 1992.

The proteins encoded by the cDNAs may also be evaluated for their influence on early steps of T-cell commitment and development. Numerous assays for such activity are familiar to those skilled in the art, including without limitation the assays disclosed in the following references, which are incorporated herein by references: Antica et al., Blood 84:111-117, 1994; Fine et al., Cell. Immunol. 155:111-122, 1994; Galy et al., Blood 85:2770-2778, 1995; Toki et al., Proc. Nat. Acad Sci. USA 88:7548-7551, 1991.

Those proteins which exhibit activity as immune system regulators activity may then be formulated as pharmaceuticals and used to treat clinical conditions in which regulation of immune activity is beneficial. For example, the protein may be useful in the treatment of various immune deficiencies and disorders (including severe combined immunodeficiency),

e.g., in regulating (up or down) growth and proliferation of T and/or B lymphocytes, as well as effecting the cytolytic activity of NK cells and other cell populations. These immune deficiencies may be genetic or be caused by viral (e.g., HIV) as well as bacterial or fungal infections, or may result from autoimmune disorders. More specifically, infectious diseases caused by viral, bacterial, fungal or other infection may be treatable using a protein encoded by extended cDNAs derived from the 5' ESTs of the present invention, including infections by HIV, hepatitis viruses, herpesviruses, mycobacteria, Leishmania spp., plamodium and various fungal infections such as candidiasis. Of course, in this regard, a protein encoded by extended cDNAs derived from the 5' ESTs of the present invention may also be useful where a boost to the immune system generally may be desirable, *i.e.*, in the treatment of cancer.

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Alternatively, proteins encoded by extended cDNAs derived from the 5' ESTs of the present invention may be used in treatment of autoimmune disorders including, for example, connective tissue disease, multiple sclerosis, systemic lupus erythematosus, rheumatoid arthritis, autoimmune pulmonary inflammation, Guillain-Barre syndrome, autoimmune thyroiditis, insulin dependent diabetes mellitis, myasthenia gravis, graft-versus-host disease and autoimmune inflammatory eye disease. Such a protein encoded by extended cDNAs derived from the 5' ESTs of the present invention may also to be useful in the treatment of allergic reactions and conditions, such as asthma (particularly allergic asthma) or other respiratory problems. Other conditions, in which immune suppression is desired (including, for example, organ transplantation), may also be treatable using a protein encoded by extended cDNAs derived from the 5' ESTs of the present invention.

Using the proteins of the invention it may also be possible to regulate immune responses either up or down.

Down regulation may involve inhibiting or blocking an immune response already in progress or may involve preventing the induction of an immune response. The functions of activated T-cells may be inhibited by suppressing T cell responses or by inducing specific tolerance in T cells, or both. Immunosuppression of T cell responses is generally an active non-antigen-specific process which requires continuous exposure of the T cells to the suppressive agent. Tolerance, which involves inducing non-responsiveness or anergy in T cells, is distinguishable from immunosuppression in that it is generally antigen-specific and persists after the end of exposure to the tolerizing agent. Operationally, tolerance can be

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demonstrated by the lack of a T cell response upon reexposure to specific antigen in the absence of the tolerizing agent.

Down regulating or preventing one or more antigen functions (including without limitation B lymphocyte antigen functions, such as, for example, B7 costimulation), e.g., preventing high level lymphokine synthesis by activated T cells, will be useful in situations of tissue, skin and organ transplantation and in graft-versus-host disease (GVHD). For example, blockage of T cell function should result in reduced tissue destruction in tissue transplantation. Typically, in tissue transplants, rejection of the transplant is initiated through its recognition as foreign by T cells, followed by an immune reaction that destroys the transplant. The administration of a molecule which inhibits or blocks interaction of a B7 lymphocyte antigen with its natural ligand(s) on immune cells (such as a soluble, monomeric form of a peptide having B7-2 activity alone or in conjunction with a monomeric form of a peptide having an activity of another B lymphocyte antigen (e.g., B7-1, B7-3) or blocking antibody), prior to transplantation, can lead to the binding of the molecule to the natural ligand(s) on the immune cells without transmitting the corresponding costimulatory signal. Blocking B lymphocyte antigen function in this matter prevents cytokine synthesis by immune cells, such as T cells, and thus acts as an immunosuppressant. Moreover, the lack of costimulation may also be sufficient to anergize the T cells, thereby inducing tolerance in a subject. Induction of long-term tolerance by B lymphocyte antigen-blocking reagents may avoid the necessity of repeated administration of these blocking reagents. To achieve sufficient immunosuppression or tolerance in a subject, it may also be necessary to block the function of a combination of B lymphocyte antigens.

The efficacy of particular blocking reagents in preventing organ transplant rejection or GVHD can be assessed using animal models that are predictive of efficacy in humans. Examples of appropriate systems which can be used include allogeneic cardiac grafts in rats and xenogeneic pancreatic islet cell grafts in mice, both of which have been used to examine the immunosuppressive effects of CTLA4Ig fusion proteins *in vivo* as described in Lenschow *et al.*, *Science* 257:789-792, 1992 and Turka *et al.*, *Proc. Natl. Acad. Sci USA*, 89:11102-11105, 1992. In addition, murine models of GVHD (see Paul ed., *Fundamental Immunology*, Raven Press, New York, 1989, pp. 846-847) can be used to determine the effect of blocking B lymphocyte antigen function *in vivo* on the development of that disease.

Blocking antigen function may also be therapeutically useful for treating autoimmune diseases. Many autoimmune disorders are the result of inappropriate activation of T cells that are reactive against self tissue and which promote the production of cytokines and autoantibodies involved in the pathology of the diseases. Preventing the activation of autoreactive T cells may reduce or eliminate disease symptoms. Administration of reagents which block costimulation of T cells by disrupting receptor/ligand interactions of B lymphocyte antigens can be used to inhibit T cell activation and prevent production of autoantibodies or T cell-derived cytokines which potentially involved in the disease process. Additionally, blocking reagents may induce antigen-specific tolerance of autoreactive T cells which could lead to long-term relief from the disease. The efficacy of blocking reagents in preventing or alleviating autoimmune disorders can be determined using a number of well-characterized animal models of human autoimmune diseases. Examples include murine experimental autoimmune encephalitis, systemic lupus erythmatosis in MRL/pr/pr mice or NZB hybrid mice, murine autoimmuno collagen arthritis, diabetes mellitus in OD mice and BB rats, and murine experimental myasthenia gravis (see Paul ed., supra, pp. 840-856).

Upregulation of an antigen function (preferably a B lymphocyte antigen function), as a means of up regulating immune responses, may also be useful in therapy. Upregulation of immune responses may involve either enhancing an existing immune response or eliciting an initial immune response as shown by the following examples. For instance, enhancing an immune response through stimulating B lymphocyte antigen function may be useful in cases of viral infection. In addition, systemic viral diseases such as influenza, the common cold, and encephalitis might be alleviated by the administration of stimulatory form of B lymphocyte antigens systemically.

Alternatively, antiviral immune responses may be enhanced in an infected patient by removing T cells from the patient, costimulating the T cells *in vitro* with viral antigen-pulsed APCs either expressing a peptide encoded by extended cDNAs derived from the 5' ESTs of the present invention or together with a stimulatory form of a soluble peptide encoded by extended cDNAs derived from the 5' ESTs of the present invention and reintroducing the *in vitro* primed T cells into the patient. The infected cells would now be capable of delivering a costimulatory signal to T cells *in vivo*, thereby activating the T cells.

In another application, upregulation or enhancement of antigen function (preferably B lymphocyte antigen function) may be useful in the induction of tumor immunity. Tumor cells (e.g., sarcoma, melanoma, lymphoma, leukemia, neuroblastoma, carcinoma) transfected with a nucleic acid encoding at least one peptide encoded by extended cDNAs derived from the 5' ESTs of the present invention can be administered to a subject to overcome tumor-specific tolerance in the subject. If desired, the tumor cell can be transfected to express a combination of peptides. For example, tumor cells obtained from a patient can be transfected *ex vivo* with an expression vector directing the expression of a peptide having B7-2-like activity alone, or in conjunction with a peptide having B7-1-like activity and/or B7-3-like activity. The transfected tumor cells are returned to the patient to result in expression of the peptides on the surface of the transfected cell. Alternatively, gene therapy techniques can be used to target a tumor cell for transfection *in vivo*.

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The presence of the peptide encoded by extended cDNAs derived from the 5' ESTs of the present invention having the activity of a B lymphocyte antigen(s) on the surface of the tumor cell provides the necessary costimulation signal to T cells to induce a T cell mediated immune response against the transfected tumor cells. In addition, tumor cells which lack or which fail to reexpress sufficient amounts of MHC class I or MHC class II molecules can be transfected with nucleic acids encoding all or a portion of (e.g., a cytoplasmic-domain truncated portion) of an MHC class I α chain and β_2 microglobulin or an MHC class II α chain and an MHC class II β chain to thereby express MHC class I or MHC class II proteins on the cell surface, respectively. Expression of the appropriate MHC class I or class II molecules in conjunction with a peptide having the activity of a B lymphocyte antigen (e.g., B7-1, B7-2, B7-3) induces a T cell mediated immune response against the transfected tumor cell. Optionally, a gene encoding an antisense construct which blocks expression of an MHC class II associated protein, such as the invariant chain, can also be cotransfected with a DNA encoding a peptide having the activity of a B lymphocyte antigen to promote presentation of tumor associated antigens and induce tumor specific immunity. Thus, the induction of a T cell mediated immune response in a human subject may be sufficient to overcome tumorspecific tolerance in the subject. Alternatively, as described in more detail below, genes encoding these immune system regulator proteins or nucleic acids regulating the expression of

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such proteins may be introduced into appropriate host cells to increase or decrease the expression of the proteins as desired.

EXAMPLE 34

Assaying the Proteins Expressed from Extended cDNAs or Portions Thereof for Hematopoiesis Regulating Activity

The proteins encoded by the extended cDNAs or portions thereof may also be evaluated for their hematopoiesis regulating activity. For example, the effect of the proteins on embryonic stem cell differentiation may be evaluated. Numerous assays for such activity are familiar to those skilled in the art, including the assays disclosed in the following references, which are incorporated herein by reference: Johansson *et al. Cell. Biol.* 15:141-151, 1995; Keller *et al.*, *Mol. Cell. Biol.* 13:473-486, 1993; McClanahan *et al.*, *Blood* 81:2903-2915, 1993.

The proteins encoded by the extended cDNAs or portions thereof may also be evaluated for their influence on the lifetime of stem cells and stem cell differentiation. Numerous assays for such activity are familiar to those skilled in the art, including the assays disclosed in the following references, which are incorporated herein by reference: Freshney, Methylcellulose Colony Forming Assays, in Culture of Hematopoietic Cells., Freshney, et al. Eds. pp. 265-268, Wiley-Liss, Inc., New York, NY. 1994; Hirayama et al., Proc. Natl. Acad. Sci. USA 89:5907-5911, 1992; McNiece and Briddell, in Culture of Hematopoietic Cells, supra, Neben et al., Exp. Hematol. 22:353-359, 1994; Ploemacher and Cobblestone In Culture of Hematopoietic Cells, supra1-21, Spooncer et al, in Culture of Hematopoietic Cells, supra 139-162.

Those proteins which exhibit hematopoiesis regulatory activity may then be formulated as pharmaceuticals and used to treat clinical conditions in which regulation of hematopoeisis is beneficial, such as in the treatment of myeloid or lymphoid cell deficiencies. Involvement in regulating hematopoiesis is indicated even by marginal biological activity in support of colony forming cells or of factor-dependent cell lines. For example, proteins supporting the growth and proliferation of erythroid progenitor cells alone or in combination with other cytokines, indicates utility, for example, in treating various anemias or for use in conjunction with irradiation/chemotherapy to stimulate the production of erythroid precursors

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and/or erythroid cells. Proteins supporting the growth and proliferation of myeloid cells such as granulocytes and monocytes/macrophages (i.e., traditional CSF activity) may be useful, for example, in conjunction with chemotherapy to prevent or treat consequent myelosuppression. Proteins supporting the growth and proliferation of megakaryocytes and consequently of platelets allows prevention or treatment of various platelet disorders such as thrombocytopenia, and generally may be used in place of or complementary to platelet transfusions. Proteins supporting the growth and proliferation of hematopoietic stem cells which are capable of maturing to any and all of the above-mentioned hematopoietic cells may therefore find therapeutic utility in various stem cell disorders (such as those usually treated with transplantion, including, without limitation, aplastic anemia and paroxysmal nocturnal hemoglobinuria), as well as in repopulating the stem cell compartment post irradiation/chemotherapy, either in vivo or ex vivo (i.e., in conjunction with bone marrow transplantation or with peripheral progenitor cell transplantation (homologous or heterologous)) as normal cells or genetically manipulated for gene therapy. Alternatively, as described in more detail below, genes encoding hematopoiesis regulating activity proteins or nucleic acids regulating the expression of such proteins may be introduced into appropriate host cells to increase or decrease the expression of the proteins as desired.

EXAMPLE 35

20 <u>Assaying the Proteins Expressed from Extended cDNAs or Portions Thereof</u> <u>for Regulation of Tissue Growth</u>

The proteins encoded by the extended cDNAs or portions thereof may also be evaluated for their effect on tissue growth. Numerous assays for such activity are familiar to those skilled in the art, including the assays disclosed in International Patent Publication No. WO95/16035, International Patent Publication No. WO95/05846 and International Patent Publication No. WO91/07491, which are incorporated herein by reference.

Assays for wound healing activity include, without limitation, those described in: Winter, *Epidermal Wound Healing*, pps. 71-112, Maibach and Rovee, eds., Year Book Medical Publishers, Inc., Chicago, as modified by Eaglstein and Mertz, *J. Invest. Dermatol.* 71:382-84, 1978, which are incorporated herein by reference.

Those proteins which are involved in the regulation of tissue growth may then be formulated as pharmaceuticals and used to treat clinical conditions in which regulation of tissue growth is beneficial. For example, a protein encoded by extended cDNAs derived from the 5' ESTs of the present invention also may have utility in compositions used for bone, cartilage, tendon, ligament and/or nerve tissue growth or regeneration, as well as for wound healing and tissue repair and replacement, and in the treatment of burns, incisions and ulcers.

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A protein encoded by extended cDNAs derived from the 5' ESTs of the present invention, which induces cartilage and/or bone growth in circumstances where bone is not normally formed, has application in the healing of bone fractures and cartilage damage or defects in humans and other animals. Such a preparation employing a protein of the invention may have prophylactic use in closed as well as open fracture reduction and also in the improved fixation of artificial joints. *De novo* bone synthesis induced by an osteogenic agent contributes to the repair of congenital, trauma induced, or oncologic resection induced craniofacial defects, and also is useful in cosmetic plastic surgery.

A protein of this invention may also be used in the treatment of periodontal disease, and in other tooth repair processes. Such agents may provide an environment to attract bone-forming cells, stimulate growth of bone-forming cells or induce differentiation of bone-forming cell progenitors. A protein of the invention may also be useful in the treatment of osteoporosis or osteoarthritis, such as through stimulation of bone and/or cartilage repair or by blocking inflammation or processes of tissue destruction (collagenase activity, osteoclast activity, etc.) mediated by inflammatory processes.

Another category of tissue regeneration activity that may be attributable to the protein encoded by extended cDNAs derived from the 5' ESTs of the present invention is tendon/ligament formation. A protein encoded by extended cDNAs derived from the 5' ESTs of the present invention, which induces tendon/ligament-like tissue or other tissue formation in circumstances where such tissue is not normally formed, has application in the healing of tendon or ligament tears, deformities and other tendon or ligament defects in humans and other animals. Such a preparation employing a tendon/ligament-like tissue inducing protein may have prophylactic use in preventing damage to tendon or ligament tissue, as well as use in the improved fixation of tendon or ligament to bone or other tissues, and in repairing defects to tendon or ligament tissue. De novo tendon/ligament-like tissue

WO 99/06550 PCT/IB98/01232

formation induced by a composition encoded by extended cDNAs derived from the 5' ESTs of the present invention contributes to the repair of tendon or ligaments defects of congenital, traumatic or other origin and is also useful in cosmetic plastic surgery for attachment or repair of tendons or ligaments. The compositions encoded by extended cDNAs derived from the 5' ESTs of the present invention may provide an environment to attract tendon- or ligament-forming cells, stimulate growth of tendon- or ligament-forming cells, induce differentiation of progenitors of tendon- or ligament-forming cells, or induce growth of tendon/ligament cells or progenitors ex vivo for return in vivo to effect tissue repair. The compositions of the invention may also be useful in the treatment of tendinitis, carpal tunnel syndrome and other tendon or ligament defects. The compositions may also include an appropriate matrix and/or sequestering agent as a carrier as is well known in the art.

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The protein encoded by extended cDNAs derived from the 5' ESTs of the present invention may also be useful for proliferation of neural cells and for regeneration of nerve and brain tissue, *i.e.*, for the treatment of central and peripheral nervous system diseases and neuropathies, as well as mechanical and traumatic disorders, which involve degeneration, death or trauma to neural cells or nerve tissue. More specifically, a protein may be used in the treatment of diseases of the peripheral nervous system, such as peripheral nerve injuries, peripheral neuropathy and localized neuropathies, and central nervous system diseases, such as Alzheimer's, Parkinson's disease, Huntington's disease, amyotrophic lateral sclerosis, and Shy-Drager syndrome. Further conditions which may be treated in accordance with the present invention include mechanical and traumatic disorders, such as spinal cord disorders, head trauma and cerebrovascular diseases such as stroke. Peripheral neuropathies resulting from chemotherapy or other medical therapies may also be treatable using a protein of the invention.

Proteins of the invention may also be useful to promote better or faster closure of non-healing wounds, including without limitation pressure ulcers, ulcers associated with vascular insufficiency, surgical and traumatic wounds, and the like.

It is expected that a protein encoded by extended cDNAs derived from the 5' ESTs of the present invention may also exhibit activity for generation or regeneration of other tissues, such as organs (including, for example, pancreas, liver, intestine, kidney, skin, endothelium) muscle (smooth, skeletal or cardiac) and vascular (including vascular

endothelium) tissue, or for promoting the growth of cells comprising such tissues. Part of the desired effects may be by inhibition or modulation of fibrotic scarring to allow normal tissue to generate. A protein of the invention may also exhibit angiogenic activity.

A protein encoded by extended cDNAs derived from the 5' ESTs of the present invention may also be useful for gut protection or regeneration and treatment of lung or liver fibrosis, reperfusion injury in various tissues, and conditions resulting from systemic cytokinc damage.

A protein encoded by extended cDNAs derived from the 5' ESTs of the present invention may also be useful for promoting or inhibiting differentiation of tissues described above from precursor tissues or cells; or for inhibiting the growth of tissues described above.

Alternatively, as described in more detail below, genes encoding tissue growth regulating activity proteins or nucleic acids regulating the expression of such proteins may be introduced into appropriate host cells to increase or decrease the expression of the proteins as desired.

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EXAMPLE 36

Assaving the Proteins Expressed from Extended cDNAs or Portions Thereof for Regulation of Reproductive Hormones

The proteins encoded by the extended cDNAs or portions thereof may also be evaluated for their ability to regulate reproductive hormones, such as follicle stimulating hormone. Numerous assays for such activity are familiar to those skilled in the art, including the assays disclosed in the following references, which are incorporated herein by reference: Vale et al., Endocrinol. 91:562-572, 1972; Ling et al., Nature 321:779-782, 1986; Vale et al., Nature 321:776-779, 1986; Mason et al., Nature 318:659-663, 1985; Forage et al., Proc. Natl. Acad. Sci. USA 83:3091-3095, 1986, Chapter 6.12 in Current Protocols in Immunology, Coligan et al. Eds. Greene Publishing Associates and Wiley-Intersciece; Taub et al., J. Clin. Invest. 95:1370-1376, 1995; Lind et al., APMIS 103:140-146, 1995; Muller et al., Eur. J. Immunol. 25:1744-1748; Gruber et al., J. Immunol. 152:5860-5867, 1994; Johnston et al., J Immunol. 153:1762-1768, 1994.

Those proteins which exhibit activity as reproductive hormones or regulators of cell movement may then be formulated as pharmaceuticals and used to treat clinical conditions in

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which regulation of reproductive hormones are beneficial. For example, a protein encoded by extended cDNAs derived from the 5' ESTs of the present invention may also exhibit activinor inhibin-related activities. Inhibins are characterized by their ability to inhibit the release of follicle stimulating hormone (FSH), while activins are characterized by their ability to stimulate the release of FSH. Thus, a protein encoded by extended cDNAs derived from the 5' ESTs of the present invention, alone or in heterodimers with a member of the inhibin α family, may be useful as a contraceptive based on the ability of inhibins to decrease fertility in female mammals and decrease spermatogenesis in male mammals. Administration of sufficient amounts of other inhibins can induce infertility in these mammals. Alternatively, the protein of the invention, as a homodimer or as a heterodimer with other protein subunits of the inhibin-B group, may be useful as a fertility inducing therapeutic, based upon the ability of activin molecules in stimulating FSH release from cells of the anterior pituitary. See, for example, United States Patent 4,798,885, the disclosure of which is incorporated herein by reference. A protein of the invention may also be useful for advancement of the onset of fertility in sexually immature mammals, so as to increase the lifetime reproductive performance of domestic animals such as cows, sheep and pigs.

Alternatively, as described in more detail below, genes encoding reproductive hormone regulating activity proteins or nucleic acids regulating the expression of such proteins may be introduced into appropriate host cells to increase or decrease the expression of the proteins as desired.

EXAMPLE 37

Assaying the Proteins Expressed from Extended cDNAs or Portions Thereof for Chemotactic/Chemokinetic Activity

The proteins encoded by the extended cDNAs or portions thereof may also be evaluated for chemotactic/chemokinetic activity. For example, a protein encoded by extended cDNAs derived from the 5' ESTs of the present invention may have chemotactic or chemokinetic activity (e.g., act as a chemokine) for mammalian cells, including, for example, monocytes, fibroblasts, neutrophils, T-cells, mast cells, eosinophils, epithelial and/or endothelial cells. Chemotactic and chemokinetic proteins can be used to mobilize or attract a desired cell population to a desired site of action. Chemotactic or chemokinetic proteins

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provide particular advantages in treatment of wounds and other trauma to tissues, as well as in treatment of localized infections. For example, attraction of lymphocytes, monocytes or neutrophils to tumors or sites of infection may result in improved immune responses against the tumor or infecting agent.

A protein or peptide has chemotactic activity for a particular cell population if it can stimulate, directly or indirectly, the directed orientation or movement of such cell population. Preferably, the protein or peptide has the ability to directly stimulate directed movement of cells. Whether a particular protein has chemotactic activity for a population of cells can be readily determined by employing such protein or peptide in any known assay for cell chemotaxis.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Assays for chemotactic activity (which will identify proteins that induce or prevent chemotaxis) consist of assays that measure the ability of a protein to induce the migration of cells across a membrane as well as the ability of a protein to induce the adhesion of one cell population to another cell population. Suitable assays for movement and adhesion include, without limitation, those described in: *Current Protocols in Immunology*, Ed by Coligan, Kruisbeek, Margulies, Shevach and Strober, Pub. Greene Publishing Associates and Wiley-Interscience, Chapter 6.12: 6.12.1-6.12.28; Taub et al., J. Clin. Invest. 95:1370-1376, 1995; Lind et al., APMIS 103:140-146, 1995; Mueller et al., Eur. J. Immunol. 25:1744-1748; Gruber et al., J. Immunol. 152:5860-5867, 1994; Johnston et al. J. Immunol., 153:1762-1768, 1994.

EXAMPLE 38

Assaying the Proteins Expressed from Extended cDNAs or

Portions Thereof for Regulation of Blood Clotting

The proteins encoded by the extended cDNAs or portions thereof may also be evaluated for their effects on blood clotting. Numerous assays for such activity are familiar to those skilled in the art, including the assays disclosed in the following references, which are incorporated herein by reference: Linet et al., J. Clin. Pharmacol. 26:131-140, 1986; Burdick

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et al., Thrombosis Res. 45:413-419, 1987; Humphrey et al., Fibrinolysis 5:71-79, 1991; Schaub, Prostaglandins 35:467-474, 1988.

Those proteins which are involved in the regulation of blood clotting may then be formulated as pharmaceuticals and used to treat clinical conditions in which regulation of blood clotting is beneficial. For example, a protein of the invention may also exhibit hemostatic or thrombolytic activity. As a result, such a protein is expected to be useful in treatment of various coagulations disorders (including hereditary disorders, such as hemophilias) or to enhance coagulation and other hemostatic events in treating wounds resulting from trauma, surgery or other causes. A protein of the invention may also be useful for dissolving or inhibiting formation of thromboses and for treatment and prevention of conditions resulting therefrom (such as infarction of cardiac and central nervous system vessels (e.g., stroke)). Alternatively, as described in more detail below, genes encoding blood clotting activity proteins or nucleic acids regulating the expression of such proteins may be introduced into appropriate host cells to increase or decrease the expression of the proteins as desired.

EXAMPLE 39

Assaying the Proteins Expressed from Extended cDNAs or Portions Thereof for Involvement in Receptor/Ligand Interactions

The proteins encoded by the extended cDNAs or a portion thereof may also be evaluated for their involvement in receptor/ligand interactions. Numerous assays for such involvement are familiar to those skilled in the art, including the assays disclosed in the following references, which are incorporated herein by reference: Chapter 7. 7.28.1-7.28.22 in Current Protocols in Immunology, Coligan et al. Eds. Greene Publishing Associates and Wiley-Interscience; Takai et al., Proc. Natl. Acad. Sci. USA 84:6864-6868, 1987; Bierer et al., J. Exp. Med. 168:1145-1156, 1988; Rosenstein et al., J. Exp. Med. 169:149-160, 1989; Stoltenborg et al., J. Immunol. Methods 175:59-68, 1994; Stitt et al., Cell 80:661-670, 1995; Gyuris et al., Cell 75:791-803, 1993.

For example, the proteins encoded by extended cDNAs derived from the 5' ESTs of the present invention may also demonstrate activity as receptors, receptor ligands or inhibitors or agonists of receptor/ligand interactions. Examples of such receptors and ligands include,

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without limitation, cytokine receptors and their ligands, receptor kinases and their ligands, receptor phosphatases and their ligands, receptors involved in cell-cell interactions and their ligands (including without limitation, cellular adhesion molecules (such as selectins, integrins and their ligands) and receptor/ligand pairs involved in antigen presentation, antigen recognition and development of cellular and humoral immune responses). Receptors and ligands are also useful for screening of potential peptide or small molecule inhibitors of the relevant receptor/ligand interaction. A protein encoded by extended cDNAs derived from the 5' ESTs of the present invention (including, without limitation, fragments of receptors and ligands) may themselves be useful as inhibitors of receptor/ligand interactions. Alternatively, as described in more detail below, genes encoding proteins involved in receptor/ligand interactions or nucleic acids regulating the expression of such proteins may be introduced into appropriate host cells to increase or decrease the expression of the proteins as desired.

EXAMPLE 40

Assaying the Proteins Expressed from Extended cDNAs or Portions Thereof for Anti-Inflammatory Activity

The proteins encoded by the extended cDNAs or a portion thereof may also be evaluated for anti-inflammatory activity. The anti-inflammatory activity may be achieved by providing a stimulus to cells involved in the inflammatory response, by inhibiting or promoting cell-cell interactions (such as, for example, cell adhesion), by inhibiting or promoting chemotaxis of cells involved in the inflammatory process, inhibiting or promoting cell extravasation, or by stimulating or suppressing production of other factors which more directly inhibit or promote an inflammatory response. Proteins exhibiting such activities can be used to treat inflammatory conditions including chronic or acute conditions, including without limitation inflammation associated with infection (such as septic shock, sepsis or systemic inflammatory response syndrome), ischemia-reperfusioninury, endotoxin lethality, arthritis, complement-mediated hyperacute rejection, nephritis, cytokine- or chemokine-induced lung injury, inflammatory bowel disease, Crohn's disease or resulting from over production of cytokines such as TNF or IL-1. Proteins of the invention may also be useful to treat anaphylaxis and hypersensitivity to an antigenic substance or material. Alternatively, as described in more detail below, genes encoding anti-inflammatory activity proteins or nucleic

acids regulating the expression of such proteins may be introduced into appropriate host cells to increase or decrease the expression of the proteins as desired.

EXAMPLE 41

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Assaying the Proteins Expressed from Extended cDNAs or Portions Thereof for Tumor Inhibition Activity

The proteins encoded by the extended cDNAs or a portion thereof may also be evaluated for tumor inhibition activity. In addition to the activities described above for immunological treatment or prevention of tumors, a protein of the invention may exhibit other anti-tumor activities. A protein may inhibit tumor growth directly or indirectly (such as, for example, via ADCC). A protein may exhibit its tumor inhibitory activity by acting on tumor tissue or tumor precursor tissue, by inhibiting formation of tissues necessary to support tumor growth (such as, for example, by inhibiting angiogenesis), by causing production of other factors, agents or cell types which inhibit tumor growth, or by suppressing, eliminating or inhibiting factors, agents or cell types which promote tumor growth. Alternatively, as described in more detail below, genes tumor inhibition activity proteins or nucleic acids regulating the expression of such proteins may be introduced into appropriate host cells to increase or decrease the expression of the proteins as desired.

A protein of the invention may also exhibit one or more of the following additional activities or effects: inhibiting the growth, infection or function of, or killing, infectious agents, including, without limitation, bacteria, viruses, fungi and other parasites; effecting (suppressing or enhancing) bodily characteristics, including, without limitation, height, weight, hair color, eye color, skin, fat to lean ratio or other tissue pigmentation, or organ or body part size or shape (such as, for example, breast augmentation or diminution, change in 25 bone form or shape); effecting biorhythms or circadian cycles or rhythms; effecting the fertility of male or female subjects; effecting the metabolism, catabolism, anabolism, processing, utilization, storage or elimination of dietary fat, lipid, protein, carbohydrate, vitamins, minerals, cofactors or other nutritional factors or component(s); effecting behavioral characteristics, including, without limitation, appetite, libido, stress, cognition (including 30

cognitive disorders), depression (including depressive disorders) and violent behaviors;

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providing analgesic effects or other pain reducing effects; promoting differentiation and growth of embryonic stem cells in lineages other than hematopoietic lineages; hormonal or endocrine activity; in the case of enzymes, correcting deficiencies of the enzyme and treating deficiency-related diseases; treatment of hyperproliferative disorders (such as, for example, psoriasis); immunoglobulin-like activity (such as, for example, the ability to bind antigens or complement); and the ability to act as an antigen in a vaccine composition to raise an immune response against such protein or another material or entity which is cross-reactive with such protein. Alternatively, as described in more detail below, genes encoding proteins involved in any of the above mentioned activities or nucleic acids regulating the expression of such proteins may be introduced into appropriate host cells to increase or decrease the expression of the proteins as desired.

EXAMPLE 42

Identification of Proteins which Interact with Polypeptides Encoded by Extended cDNAs

Proteins which interact with the polypeptides encoded by cDNAs derived from the 5' ESTs or fragments thereof, such as receptor proteins, may be identified using two hybrid systems such as the Matchmaker Two Hybrid System 2 (Catalog No. K1604-1, Clontech). As described in the manual accompanying the kit which is incorporated herein by reference, the the cDNAs derived from 5' ESTs, or fragments thereof, are inserted into an expression vector such that they are in frame with DNA encoding the DNA binding domain of the yeast

encoded by the extended cDNAs or portions thereof.

wector such that they are in frame with DNA encoding the DNA binding domain of the yeast transcriptional activator GAL4. cDNAs in a cDNA library which encode proteins which might interact with the polypeptides encoded by the extended cDNAs or portions thereof are inserted into a second expression vector such that they are in frame with DNA encoding the activation domain of GAL4. The two expression plasmids are transformed into yeast and the yeast are plated on selection medium which selects for expression of selectable markers on each of the expression vectors as well as GAL4 dependent expression of the HIS3 gene. Transformants capable of growing on medium lacking histidine are screened for GAL4 dependent lacZ expression. Those cells which are positive in both the histidine selection and the lacZ assay contain plasmids encoding proteins which interact with the polypeptide

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Alternatively, the system described in Lustig et al., Methods in Enzymology 283: 83-99, 1997, and in U.S. Patent No. 5,654,150, the disclosure of which is incorporated herein by reference, may be used for identifying molecules which interact with the polypeptides encoded by extended cDNAs. In such systems, in vitro transcription reactions are performed on a pool of vectors containing extended cDNA inserts cloned downstream of a promoter which drives in vitro transcription. The resulting pools of mRNAs are introduced into Xenopus laevis oocytes. The oocytes are then assayed for a desired activity.

Alternatively, the pooled *in vitro* transcription products produced as described above may be translated *in vitro*. The pooled *in vitro* translation products can be assayed for a desired activity or for interaction with a known polypeptide.

Proteins or other molecules interacting with polypeptides encoded by extended cDNAs can be found by a variety of additional techniques. In one method, affinity columns containing the polypeptide encoded by the extended cDNA or a portion thereof can be constructed. In some versions, of this method the affinity column contains chimeric proteins in which the protein encoded by the extended cDNA or a portion thereof is fused to glutathione S-transferase. A mixture of cellular proteins or pool of expressed proteins as described above and is applied to the affinity column. Proteins interacting with the polypeptide attached to the column can then be isolated and analyzed on 2-D electrophoresis gel as described in Ramunsen *et al.*, *Electrophoresis* 18:588-598, 1997, the disclosure of which is incorporated herein by reference. Alternatively, the proteins retained on the affinity column can be purified by electrophoresis based methods and sequenced. The same method can be used to isolate antibodies, to screen phage display products, or to screen phage display human antibodies.

Proteins interacting with polypeptides encoded by extended cDNAs or portions thereof can also be screened by using an Optical Biosensor as described in Edwards and Leatherbarrow, Analytical Biochemistry 246:1-6, 1997, the disclosure of which is incorporated herein by reference. The main advantage of the method is that it allows the determination of the association rate between the protein and other interacting molecules. Thus, it is possible to specifically select interacting molecules with a high or low association rate. Typically a target molecule is linked to the sensor surface (through a carboxymethl dextran matrix) and a sample of test molecules is placed in contact with

WO 99/06550 PCT/IB98/01232

the target molecules. The binding of a test molecule to the target molecule causes a change in the refractive index and/ or thickness. This change is detected by the Biosensor provided it occurs in the evanescent field (which extend a few hundred nanometers from the sensor surface). In these screening assays, the target molecule can be one of the polypeptides encoded by extended cDNAs or a portion thereof and the test sample can be a collection of proteins extracted from tissues or cells, a pool of expressed proteins, combinatorial peptide and/ or chemical libraries, or phage displayed peptides. The tissues or cells from which the test proteins are extracted can originate from any species.

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In other methods, a target protein is immobilized and the test population is a collection of unique polypeptides encoded by the extended cDNAs or portions thereof.

To study the interaction of the proteins encoded by the extended cDNAs or portions thereof with drugs, the microdialysis coupled to HPLC method described by Wang et al., Chromatographia 44:205-208, 1997 or the affinity capillary electrophoresis method described by Busch et al., J. Chromatogr. 777:311-328, 1997, the disclosures of which are incorporated herein by reference can be used.

It will be appreciated by those skilled in the art that the proteins expressed from the extended cDNAs or portions may be assayed for numerous activities in addition to those specifically enumerated above. For example, the expressed proteins may be evaluated for applications involving control and regulation of inflammation, tumor proliferation or metastasis, infection, or other clinical conditions. In addition, the proteins expressed from the extended cDNAs or portions thereof may be useful as nutritional agents or cosmetic agents.

The proteins expressed from the cDNAs or portions thereof may be used to generate antibodies capable of specifically binding to the expressed protein or fragments thereof as described in Example 40 below. The antibodies may capable of binding a full length protein encoded by a cDNA derived from a 5' EST, a mature protein (i.e. the protein generated by cleavage of the signal peptide) encoded by a cDNA derived from a 5' EST, or a signal peptide encoded by a cDNA derived from a 5' EST. Alternatively, the antibodies may be capable of binding fragments of at least 10 amino acids of the proteins encoded by the above cDNAs. In some embodiments, the antibodies may be capable of binding fragments of at

least 15 amino acids of the proteins encoded by the above cDNAs. In other embodiments, the antibodies may be capable of binding fragments of at least 25 amino acids of the proteins expressed from the extended cDNAs which comprise at least 25 amino acids of the proteins encoded by the above cDNAs. In further embodiments, the antibodies may be capable of binding fragments of at least 40 amino acids of the proteins encoded by the above cDNAs.

EXAMPLE 43

Production of an Antibody to a Human Protein

Substantially pure protein or polypeptide is isolated from the transfected or transformed cells as described in Example 30. The concentration of protein in the final preparation is adjusted, for example, by concentration on an Amicon filter device, to the level of a few µg/ml. Monoclonal or polyclonal antibody to the protein can then be prepared as follows:

15 1. Monoclonal Antibody Production by Hybridoma Fusion

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Monoclonal antibody to epitopes of any of the peptides identified and isolated as described can be prepared from murine hybridomas according to the classical method of Kohler, and Milstein, Nature 256:495, 1975 or derivative methods thereof. Briefly, a mouse is repetitively inoculated with a few micrograms of the selected protein or peptides derived therefrom over a period of a few weeks. The mouse is then sacrificed, and the antibody producing cells of the spleen isolated. The spleen cells are fused by means of polyethylene glycol with mouse myeloma cells, and the excess unfused cells destroyed by growth of the system on selective media comprising aminopterin (HAT media). The successfully fused cells are diluted and aliquots of the dilution placed in wells of a microtiter plate where growth of the culture is continued. Antibody-producing clones are identified by detection of antibody in the supernatant fluid of the wells by immunoassay procedures, such as ELISA, as originally described by Engvall, Meth. Enzymol. 70:419, 1980, the disclosure of which is incorporated herein by reference and derivative methods thereof. Selected positive clones can be expanded and their monoclonal antibody product harvested for use. Detailed procedures for monoclonal antibody production are described in Davis et al. in Basic Methods in Molecular Biology

WO 99/06550 PCT/IB98/01232

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Elsevier, New York. Section 21-2, the disclosure of which is incorporated herein by reference.

2. Polyclonal Antibody Production by Immunization

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Polyclonal antiserum containing antibodies to heterogenous epitopes of a single protein can be prepared by immunizing suitable animals with the expressed protein or peptides derived therefrom, which can be unmodified or modified to enhance immunogenicity. Effective polyclonal antibody production is affected by many factors related both to the antigen and the host species. For example, small molecules tend to be less immunogenic than others and may require the use of carriers and adjuvant. Also, host animals response vary depending on site of inoculations and doses, with both inadequate or excessive doses of antigen resulting in low titer antisera. Small doses (ng level) of antigen administered at multiple intradermal sites appears to be most reliable. An effective immunization protocol for rabbits can be found in Vaitukaitis. et al, J. Clin. Endocrinol. Metab. 33:988-991 (1971), the disclosure of which is incorporated herein by reference.

Booster injections can be given at regular intervals, and antiserum harvested when antibody titer thereof, as determined semi-quantitatively, for example, by double immunodiffusion in agar against known concentrations of the antigen, begins to fall. See, for example, Ouchterlony, et al., Chap. 19 in: Handbook of Experimental Immunology D. Wier (ed) Blackwell (1973), the disclosure of which is incorporated herein by reference. Plateau concentration of antibody is usually in the range of 0.1 to 0.2 mg/ml of serum (about 12 µM). Affinity of the antisera for the antigen is determined by preparing competitive binding curves, as described, for example, by Fisher, D., Chap. 42 in: Manual of Clinical Immunology, 2d Ed. (Rose and Friedman, Eds.) Amer. Soc. For Microbiol., Washington, D.C. (1980), the disclosure of which is incorporated herein by reference.

Antibody preparations prepared according to either protocol are useful in quantitative immunoassays which determine concentrations of antigen-bearing substances in biological samples; they are also used semi-quantitatively or qualitatively to identify the presence of antigen in a biological sample. The antibodies may also be used in therapeutic compositions for killing cells expressing the protein or reducing the levels of the protein in the body.

V. Use of 5' ESTs or Sequences Obtainable Therefrom or Portions Thereof as Reagents

The 5' ESTs of the present invention (or cDNAs or genomic DNAs obtainable therefrom) may be used as reagents in isolation procedures, diagnostic assays, and forensic procedures. For example, sequences from the 5' ESTs (or cDNAs or genomic DNAs obtainable therefrom) may be detectably labeled and used as probes to isolate other sequences capable of hybridizing to them. In addition, sequences from 5' ESTs (or cDNAs or genomic DNAs obtainable therefrom) may be used to design PCR primers to be used in isolation, diagnostic, or forensic procedures.

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1. Use of 5' ESTs or Sequences Obtainable Therefrom or Portions Thereof in Isolation, Diagnostic and Forensic Procedures

EXAMPLE 44

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Preparation of PCR Primers and Amplification of DNA

The 5' EST sequences (or cDNAs or genomic DNAs obtainable therefrom) may be used to prepare PCR primers for a variety of applications, including isolation procedures for cloning nucleic acids capable of hybridizing to such sequences, diagnostic techniques and forensic techniques. The PCR primers are at least 10 bases, and preferably at least 12, 15, or 17 bases in length. More preferably, the PCR primers are at least 20-30 bases in length. In some embodiments, the PCR primers may be more than 30 bases in length. It is preferred that the primer pairs have approximately the same G/C ratio, so that melting temperatures are approximately the same. A variety of PCR techniques are familiar to those skilled in the art. For a review of PCR technology, see Molecular Cloning to Genetic Engineering, White Ed. in Methods in Molecular Biology 67: Humana Press, Totowa 1997, the disclosure of which is incorporated herein by reference. In each of these PCR procedures, PCR primers on either side of the nucleic acid sequences to be amplified are added to a suitably prepared nucleic acid sample along with dNTPs and a thermostable polymerase such as Taq polymerase, Pfu polymerase, or Vent polymerase. The nucleic acid in the sample is denatured and the PCR primers are specifically hybridized to complementary nucleic acid sequences in the sample. The hybridized primers are extended. Thereafter, another cycle of denaturation,

hybridization, and extension is initiated. The cycles are repeated multiple times to produce an amplified fragment containing the nucleic acid sequence between the primer sites.

EXAMPLE 45

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Use of 5'ESTs as Probes

Probes derived from 5' ESTs (or cDNAs or genomic DNAs obtainable therefrom), including full length cDNAs or genomic sequences, may be labeled with detectable labels familiar to those skilled in the art, including radioisotopes and non-radioactive labels, to provide a detectable probe. The detectable probe may be single stranded or double stranded and may be made using techniques known in the art, including *in vitro* transcription, nick translation, or kinase reactions. A nucleic acid sample containing a sequence capable of hybridizing to the labeled probe is contacted with the labeled probe. If the nucleic acid in the sample is double stranded, it may be denatured prior to contacting the probe. In some applications, the nucleic acid sample may be immobilized on a surface such as a nitrocellulose or nylon membrane. The nucleic acid sample may comprise nucleic acids obtained from a variety of sources, including genomic DNA, cDNA libraries, RNA, or tissue samples.

Procedures used to detect the presence of nucleic acids capable of hybridizing to the detectable probe include well known techniques such as Southern blotting, Northern blotting, dot blotting, colony hybridization, and plaque hybridization. In some applications, the nucleic acid capable of hybridizing to the labeled probe may be cloned into vectors such as expression vectors, sequencing vectors, or *in vitro* transcription vectors to facilitate the characterization and expression of the hybridizing nucleic acids in the sample. For example, such techniques may be used to isolate and clone sequences in a genomic library or cDNA library which are capable of hybridizing to the detectable probe as described in Example 30 above.

PCR primers made as described in Example 44 above may be used in forensic analyses, such as the DNA fingerprinting techniques described in Examples 46-50 below. Such analyses may utilize detectable probes or primers based on the sequences of the the 5' ESTs or of cDNAs or genomic DNAs isolated using the 5' ESTs.

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EXAMPLE 46

In one exemplary method, DNA samples are isolated from forensic specimens of, for example, hair, semen, blood or skin cells by conventional methods. A panel of PCR primers based on a number of the 5' ESTs of Example 25, or cDNAs or genomic DNAs isolated therefrom as described above, is then utilized in accordance with Example 44 to amplify DNA of approximately 100-200 bases in length from the forensic specimen. Corresponding sequences are obtained from a test subject. Each of these identification DNAs is then sequenced using standard techniques, and a simple database comparison determines the differences, if any, between the sequences from the subject and those from the sample. Statistically significant differences between the suspect's DNA sequences and those from the sample conclusively prove a lack of identity. This lack of identity can be proven, for example, with only one sequence. Identity, on the other hand, should be demonstrated with a large number of sequences, all matching. Preferably, a minimum of 50 statistically identical sequences of 100 bases in length are used to prove identity between the suspect and the sample.

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EXAMPLE 47

Positive Identification by DNA Sequencing

The technique outlined in the previous example may also be used on a larger scale to provide a unique fingerprint-type identification of any individual. In this technique, primers are prepared from a large number of 5'EST sequences from Example 25, or cDNA or genomic DNA sequences obtainable therefrom. Preferably, 20 to 50 different primers are used. These primers are used to obtain a corresponding number of PCR-generated DNA segments from the individual in question in accordance with Example 44. Each of these DNA segments is sequenced, using the methods set forth in Example 46. The database of sequences generated through this procedure uniquely identifies the individual from whom the sequences were obtained. The same panel of primers may then be used at any later time to absolutely correlate tissue or other biological specimen with that individual.

EXAMPLE 48

WO 99/06550 PCT/IB98/01232

The procedure of Example 47 is repeated to obtain a panel of at least 10 amplified sequences from an individual and a specimen. Preferably, the panel contains at least 50 amplified sequences. More preferably, the panel contains 100 amplified sequences. In some embodiments, the panel contains 200 amplified sequences. This PCR-generated DNA is then digested with one or a combination of, preferably, four base specific restriction enzymes. Such enzymes are commercially available and known to those of skill in the art. After digestion, the resultant gene fragments are size separated in multiple duplicate wells on an agarose gel and transferred to nitrocellulose using Southern blotting techniques well known to those with skill in the art. For a review of Southern blotting see Davis *et al.* (Basic Methods in Molecular Biology, 1986, Elsevier Press. pp 62-65), the disclosure of which is incorporated herein by reference.

A panel of probes based on the sequences of 5' ESTs (or cDNAs or genomic DNAs obtainable therefrom), or fragments thereof of at least 10 bases, are radioactively or colorimetrically labeled using methods known in the art, such as nick translation or end labeling, and hybridized to the Southern blot using techniques known in the art (Davis *et al.*, supra). Preferably, the probe comprises at least 12, 15, or 17 consecutive nucleotides from the 5' EST (or cDNAs or genomic DNAs obtainable therefrom). More preferably, the probe comprises at least 20-30 consecutive nucleotides from the 5' EST (or cDNAs or genomic DNAs obtainable therefrom). In some embodiments, the probe comprises more than 30 nucleotides from the 5' EST (or cDNAs or genomic DNAs obtainable therefrom).

Preferably, at least 5 to 10 of these labeled probes are used, and more preferably at least about 20 or 30 are used to provide a unique pattern. The resultant bands appearing from the hybridization of a large sample of 5' EST (or cDNAs or genomic DNAs obtainable therefrom) will be a unique identifier. Since the restriction enzyme cleavage will be different for every individual, the band pattern on the Southern blot will also be unique. Increasing the number of 5' EST (or cDNAs or genomic DNAs obtainable therefrom) probes will provide a statistically higher level of confidence in the identification since there will be an increased number of sets of bands used for identification.

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EXAMPLE 49

Dot Blot Identification Procedure

Another technique for identifying individuals using the 5' EST sequences disclosed herein utilizes a dot blot hybridization technique.

Genomic DNA is isolated from nuclei of subject to be identified. Oligonucleotide probes of approximately 30 bp in length are synthesized that correspond to at least 10, preferably 50 sequences from the 5' ESTs or cDNAs or genomic DNAs obtainable therefrom. The probes are used to hybridize to the genomic DNA through conditions known to those in the art. The oligonucleotides are end labeled with P³² using polynucleotide kinase (Pharmacia). Dot Blots are created by spotting the genomic DNA onto nitrocellulose or the like using a vacuum dot blot manifold (BioRad, Richmond California). The nitrocellulose filter containing the genomic sequences is baked or UV linked to the filter, prehybridized and hybridized with labeled probe using techniques known in the art (Davis et al., supra). The ³²P labeled DNA fragments are sequentially hybridized with successively stringent conditions to detect minimal differences between the 30 bp sequence and the DNA. Tetramethylammonium chloride is useful for identifying clones containing small numbers of nucleotide mismatches (Wood et al., Proc. Natl. Acad. Sci. USA 82(6):1585-1588, 1985) which is hereby incorporated by reference. A unique pattern of dots distinguishes one individual from another individual.

5' EST sequences (or cDNAs or genomic DNAs obtainable therefrom) or oligonucleotides containing at least 10 consecutive bases from these sequences can be used as probes in the following alternative fingerprinting technique. Preferably, the probe comprises at least 12, 15, or 17 consecutive nucleotides from the 5' EST sequences (or cDNAs or genomic DNAs obtainable therefrom). More preferably, the probe comprises at least 20-30 consecutive nucleotides from the 5' EST sequences (or cDNAs or genomic DNAs obtainable therefrom). In some embodiments, the probe comprises more than 30 nucleotides from the 5' EST sequences (or cDNAs or genomic DNAs obtainable therefrom).

Preferably, a plurality of probes having sequences from different genes are used in the alternative fingerprinting technique. Example 50 below provides a representative alternative fingerprinting procedure in which the probes are derived from 5'EST.

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EXAMPLE 50

Alternative "Fingerprint" Identification Technique

20-mer oligonucleotides are prepared from a large number, e.g. 50, 100, or 200, of 5'EST using commercially available oligonucleotide services such as Genset, Paris, France. Cell samples from the test subject are processed for DNA using techniques well known to those with skill in the art. The nucleic acid is digested with restriction enzymes such as EcoRI and XbaI. Following digestion, samples are applied to wells for electrophoresis. The procedure, as known in the art, may be modified to accommodate polyacrylamide electrophoresis, however in this example, samples containing 5 ug of DNA are loaded into wells and separated on 0.8% agarose gels. The gels are transferred onto nitrocellulose using standard Southern blotting techniques.

10 ng of each of the oligonucleotides are pooled and end-labeled with ³²P. The nitrocellulose is prehybridized with blocking solution and hybridized with the labeled probes. Following hybridization and washing, the nitrocellulose filter is exposed to X-Omat AR X-ray film. The resulting hybridization pattern will be unique for each individual.

It is additionally contemplated within this example that the number of probe sequences used can be varied for additional accuracy or clarity.

The proteins encoded by the extended cDNAs may also be used to generate antibodies as explained in Examples 30 and 43 in order to identify the tissue type or cell species from which a sample is derived as described in example 51.

EXAMPLE 51

Identification of Tissue Types or Cell Species by Means of

Labeled Tissue Specific Antibodies

Identification of specific tissues is accomplished by the visualization of tissue specific antigens by means of antibody preparations according to Examples 30 and 43 which are conjugated, directly or indirectly to a detectable marker. Selected labeled antibody species bind to their specific antigen binding partner in tissue sections, cell suspensions, or in extracts of soluble proteins from a tissue sample to provide a pattern for qualitative or semi-qualitative interpretation.

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Antisera for these procedures must have a potency exceeding that of the native preparation, and for that reason, antibodies are concentrated to a mg/ml level by isolation of the gamma globulin fraction, for example, by ion-exchange chromatography or by ammonium sulfate fractionation. Also, to provide the most specific antisera, unwanted antibodies, for example to common proteins, must be removed from the gamma globulin fraction, for example by means of insoluble immunoabsorbents, before the antibodies are labeled with the marker. Either monoclonal or heterologous antisera is suitable for either procedure.

A. Immunohistochemical techniques

Purified, high-titer antibodies, prepared as described above, are conjugated to a detectable marker, as described, for example, by Fudenberg, Chap. 26 in: Basic and Clinical Immunology, 3rd Ed. Lange, Los Altos, California, 1980, or Rose, et al., Chap. 12 in: Methods in Immunodiagnosis, 2d Ed. John Wiley and Sons, New York (1980), the disclosures of which are incorporated herein by reference.

A fluorescent marker, either fluorescein or rhodamine, is preferred, but antibodies can also be labeled with an enzyme that supports a color producing reaction with a substrate, such as horseradish peroxidase. Markers can be added to tissue-bound antibody in a second step, as described below. Alternatively, the specific antitissue antibodies can be labeled with ferritin or other electron dense particles, and localization of the ferritin coupled antigen-antibody complexes achieved by means of an electron microscope. In yet another approach, the antibodies are radiolabeled, with, for example ¹²⁵I, and detected by overlaying the antibody treated preparation with photographic emulsion.

Preparations to carry out the procedures can comprise monoclonal or polyclonal antibodies to a single protein or peptide identified as specific to a tissue type, for example, brain tissue, or antibody preparations to several antigenically distinct tissue specific antigens can be used in panels, independently or in mixtures, as required.

Tissue sections and cell suspensions are prepared for immunohistochemical examination according to common histological techniques. Multiple cryostat sections (about 4 μ m, unfixed) of the unknown tissue and known control, are mounted and each slide covered with different dilutions of the antibody preparation. Sections of known and unknown tissues should also be treated with preparations to provide a positive control, a negative

control, for example, pre-immune sera, and a control for non-specific staining, for example, buffer.

Treated sections are incubated in a humid chamber for 30 min at room temperature, rinsed, then washed in buffer for 30-45 min. Excess fluid is blotted away, and the marker developed.

If the tissue specific antibody was not labeled in the first incubation, it can be labeled at this time in a second antibody-antibody reaction, for example, by adding fluorescein- or enzyme-conjugated antibody against the immunoglobulin class of the antiserum-producing species, for example, fluorescein labeled antibody to mouse IgG. Such labeled sera are commercially available.

The antigen found in the tissues by the above procedure can be quantified by measuring the intensity of color or fluorescence on the tissue section, and calibrating that signal using appropriate standards.

B. Identification of tissue specific soluble proteins

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The visualization of tissue specific proteins and identification of unknown tissues from that procedure is carried out using the labeled antibody reagents and detection strategy as described for immunohistochemistry; however the sample is prepared according to an electrophoretic technique to distribute the proteins extracted from the tissue in an orderly array on the basis of molecular weight for detection.

A tissue sample is homogenized using a Virtis apparatus; cell suspensions are disrupted by Dounce homogenization or osmotic lysis, using detergents in either case as required to disrupt cell membranes, as is the practice in the art. Insoluble cell components such as nuclei, microsomes, and membrane fragments are removed by ultracentrifugation, and the soluble protein-containing fraction concentrated if necessary and reserved for analysis.

A sample of the soluble protein solution is resolved into individual protein species by conventional SDS polyacrylamide electrophoresis as described, for example, by Davis, et al., Section 19-2 in: Basic Methods in Molecular Biology, Leder ed., Elsevier, New York, 1986, the disclosure of which is incorporated herein by reference, using a range of amounts of polyacrylamide in a set of gels to resolve the entire molecular weight range of proteins to be detected in the sample. A size marker is run in parallel for purposes of estimating molecular weights of the constituent proteins. Sample size for analysis is a convenient volume of from 5

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to 55 µl, and containing from about 1 to 100 µg protein. An aliquot of each of the resolved proteins is transferred by blotting to a nitrocellulose filter paper, a process that maintains the pattern of resolution. Multiple copies are prepared. The procedure, known as Western Blot Analysis, is well described in Davis, L. et al., supra Section 19-3. One set of nitrocellulose blots is stained with Coomassie blue dye to visualize the entire set of proteins for comparison with the antibody bound proteins. The remaining nitrocellulose filters are then incubated with a solution of one or more specific antisera to tissue specific proteins prepared as described in Examples 30 and 43. In this procedure, as in procedure A above, appropriate positive and negative sample and reagent controls are run.

In either procedure A or B, a detectable label can be attached to the primary tissue antigen-primary antibody complex according to various strategies and permutations thereof. In a straightforward approach, the primary specific antibody can be labeled; alternatively, the unlabeled complex can be bound by a labeled secondary anti-IgG antibody. In other approaches, either the primary or secondary antibody is conjugated to a biotin molecule, which can, in a subsequent step, bind an avidin conjugated marker. According to yet another strategy, enzyme labeled or radioactive protein A, which has the property of binding to any IgG, is bound in a final step to either the primary or secondary antibody.

The visualization of tissue specific antigen binding at levels above those seen in control tissues to one or more tissue specific antibodies, prepared from the gene sequences identified from extended cDNA sequences, can identify tissues of unknown origin, for example, forensic samples, or differentiated tumor tissue that has metastasized to foreign bodily sites.

In addition to their applications in forensics and identification, 5' ESTs (or cDNAs or genomic DNAs obtainable therefrom) may be mapped to their chromosomal locations. Example 52 below describes radiation hybrid (RH) mapping of human chromosomal regions using 5'ESTs. Example 53 below describes a representative procedure for mapping an 5' EST to its location on a human chromosome. Example 54 below describes mapping of 5' ESTs on metaphase chromosomes by Fluorescence In Situ Hybridization (FISH). Those skilled in the art will appreciate that the method of Examples 52-54 may also be used to map cDNAs or genomic DNAs obtainable from the 5' ESTs to their chromosomal locations.

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2. Use of 5' ESTs or Sequences Obtainable Therefrom or Portions Thereof in Chromosome Mapping

EXAMPLE 52

Radiation hybrid mapping of 5'ESTs to the human genome

Radiation hybrid (RH) mapping is a somatic cell genetic approach that can be used for high resolution mapping of the human genome. In this approach, cell lines containing one or more human chromosomes are lethally irradiated, breaking each chromosome into fragments whose size depends on the radiation dose. These fragments are rescued by fusion with cultured rodent cells, yielding subclones containing different portions of the human genome. This technique is described by Benham et al., Genomics 4:509-517, 1989; and Cox et al., Science 250:245-250, 1990, the entire contents of which are hereby incorporated by reference. The random and independent nature of the subclones permits efficient mapping of any human genome marker. Human DNA isolated from a panel of 80-100 cell lines provides a mapping reagent for ordering 5'EST. In this approach, the frequency of breakage between markers is used to measure distance, allowing construction of fine resolution maps as has been done using conventional ESTs (Schuler et al., Science 274:540-546, 1996, hereby incorporated by reference).

RH mapping has been used to generate a high-resolution whole genome radiation hybrid map of human chromosome 17q22-q25.3 across the genes for growth hormone (GH) and thymidine kinase (TK) (Foster et al., Genomics 33:185-192, 1996), the region surrounding the Gorlin syndrome gene (Obermayr et al., Eur. J. Hum. Genet. 4:242-245, 1996), 60 loci covering the entire short arm of chromosome 12 (Raeymaekers et al., Genomics 29:170-178, 1995), the region of human chromosome 22 containing the neurofibromatosis type 2 locus (Frazer et al., Genomics 14:574-584, 1992) and 13 loci on the long arm of chromosome 5 (Warrington et al., Genomics 11:701-708, 1991).

EXAMPLE 53

Mapping of 5'ESTs to HumanChromosomes using PCR techniques

5' ESTs (or cDNAs or genomic DNAs obtainable therefrom) may be assigned to human chromosomes using PCR based methodologies. In such approaches, oligonucleotide primer pairs are designed from the 5' ESTs (or cDNAs or genomic DNAs obtainable

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therefrom) to minimize the chance of amplifying through an intron. Preferably, the oligonucleotide primers are 18-23 bp in length and are designed for PCR amplification. The creation of PCR primers from known sequences is well known to those with skill in the art. For a review of PCR technology see Erlich in PCR Technology, Principles and Applications for DNA Amplification, Freeman and Co., New York, 1992, the disclosure of which is incorporated herein by reference.

The primers are used in polymerase chain reactions (PCR) to amplify templates from total human genomic DNA. PCR conditions are as follows: 60 ng of genomic DNA is used as a template for PCR with 80 ng of each oligonucleotide primer, 0.6 unit of Taq polymerase, and 1 μCu of a ³²P-labeled deoxycytidine triphosphate. The PCR is performed in a microplate thermocycler (Techne) under the following conditions: 30 cycles of 94°C, 1.4 min; 55°C, 2 min; and 72°C, 2 min; with a final extension at 72°C for 10 min. The amplified products are analyzed on a 6% polyacrylamide sequencing gel and visualized by autoradiography. If the length of the resulting PCR product is identical to the distance between the ends of the primer sequences in the extended cDNA from which the primers are derived, then the PCR reaction is repeated with DNA templates from two panels of human-rodent somatic cell hybrids, BIOS PCRable DNA (BIOS Corporation) and NIGMS Human-Rodent Somatic Cell Hybrid Mapping Panel Number 1 (NIGMS, Camden, NJ).

PCR is used to screen a series of somatic cell hybrid cell lines containing defined sets of human chromosomes for the presence of a given 5' EST (or cDNA or genomic DNA obtainable therefrom). DNA is isolated from the somatic hybrids and used as starting templates for PCR_reactions using the primer pairs from the 5' EST (or cDNA or genomic DNA obtainable therefrom). Only those somatic cell hybrids with chromosomes containing the human gene corresponding to the 5' EST (or cDNA or genomic DNA obtainable therefrom) will yield an amplified fragment. The 5' EST (or cDNA or genomic DNA obtainable therefrom) are assigned to a chromosome by analysis of the segregation pattern of PCR products from the somatic hybrid DNA templates. The single human chromosome present in all cell hybrids that give rise to an amplified fragment is the chromosome containing that 5'EST (or cDNA or genomic DNA obtainable therefrom). For a review of techniques and analysis of results from somatic cell gene mapping experiments, see Ledbetter *et al.*, *Genomics* 6:475-481, 1990, the disclosure of which is incorporated herein by reference.

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EXAMPLE 54

Mapping of Extended 5' ESTs to Chromosomes Using Fluorescence In Situ Hybridization

Fluorescence in situ hybridization allows the 5'EST (or cDNA or genomic DNA obtainable therefrom) to be mapped to a particular location on a given chromosome. The chromosomes to be used for fluorescence in situ hybridization techniques may be obtained from a variety of sources including cell cultures, tissues, or whole blood.

In a preferred embodiment, chromosomal localization of an 5'EST (or cDNA or genomic DNA obtainable therefrom) is obtained by FISH as described by Cherif et al. (Proc. Natl. Acad. Sci. U.S.A., 87:6639-6643, 1990), the disclosure of which is incorporated herein by reference.. Metaphase chromosomes are prepared from phytohemagglutinin (PHA)stimulated blood cell donors. PHA-stimulated lymphocytes from healthy males are cultured for 72 h in RPMI-1640 medium. For synchronization, methotrexate (10 μM) is added for 17 h, followed by addition of 5-bromodeoxyuridine (5-BrdU, 0.1 mM) for 6 h. Colcemid (1 $\mu g/ml$) is added for the last 15 min before harvesting the cells. Cells are collected, washed in RPMI, incubated with a hypotonic solution of KCl (75 mM) at 37°C for 15 min and fixed in three changes of methanol:acetic acid (3:1). The cell suspension is dropped onto a glass slide and air dried. The 5'EST (or cDNA or genomic DNA obtainable therefrom) is labeled with biotin-16 dUTP by nick translation according to the manufacturer's instructions (Bethesda Research Laboratories, Bethesda, MD), purified using a Sephadex G-50 column (Pharmacia, Upsala, Sweden) and precipitated. Just prior to hybridization, the DNA pellet is dissolved in hybridization buffer (50% formamide, 2 X SSC, 10% dextran sulfate, 1 mg/ml sonicated salmon sperm DNA, pH 7) and the probe is denatured at 70°C for 5-10 min.

Slides kept at -20°C are treated for 1 h at 37°C with RNase A (100 µg/ml), rinsed three times in 2 X SSC and dehydrated in an ethanol series. Chromosome preparations are denatured in 70% formamide, 2 X SSC for 2 min at 70°C, then dehydrated at 4°C. The slides are treated with proteinase K (10 µg/100 ml in 20 mM Tris-HCl, 2 mM CaCl₂) at 37°C for 8 min and dehydrated. The hybridization mixture containing the probe is placed on the slide, covered with a coverslip, sealed with rubber cement and incubated overnight in a humid chamber at 37°C. After hybridization and post-hybridization washes, the biotinylated probe is detected by avidin-FITC and amplified with additional layers of biotinylated goat anti-avidin

and avidin-FITC. For chromosomal localization, fluorescent R-bands are obtained as previously described (Cherif et al., supra.). The slides are observed under a LEICA fluorescence microscope (DMRXA). Chromosomes are counterstained with propidium iodide and the fluorescent signal of the probe appears as two symmetrical yellow-green spots on both chromatids of the fluorescent R-band chromosome (red). Thus, a particular 5'EST (or cDNA or genomic DNA obtainable therefrom) may be localized to a particular cytogenetic R-band on a given chromosome.

Once the 5'EST (or cDNA or genomic DNA obtainable therefrom) have been assigned to particular chromosomes using the techniques described in Examples 52-54 above, they may be utilized to construct a high resolution map of the chromosomes on which they are located or to identify the chromosomes in a sample.

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Use of 5'EST to Construct or Expand Chromosome Maps

Chromosome mapping involves assigning a given unique sequence to a particular chromosome as described above. Once the unique sequence has been mapped to a given chromosome, it is ordered relative to other unique sequences located on the same chromosome. One approach to chromosome mapping utilizes a series of yeast artificial chromosomes (YACs) bearing several thousand long inserts derived from the chromosomes of the organism from which the extended cDNAs (or genomic DNAs obtainable therefrom) are obtained. This approach is described in Nagaraja et al., Genome Research 7:210-222, 1997, the disclosure of which is incorporated herein by reference. Briefly, in this approach each chromosome is broken into overlapping pieces which are inserted into the YAC vector. The YAC inserts are screened using PCR or other methods to determine whether they include the 5'EST (or cDNA or genomic DNA obtainable therefrom) whose position is to be determined. Once an insert has been found which includes the 5'EST (or cDNA or genomic DNA obtainable therefrom), the insert can be analyzed by PCR or other methods to determine whether the insert also contains other sequences known to be on the chromosome or in the region from which the 5'EST (or cDNA or genomic DNA obtainable therefrom) was derived. This process can be repeated for each insert in the YAC library to determine the

location of each of the extended cDNAs (or genomic DNAs obtainable therefrom) relative to one another and to other known chromosomal markers. In this way, a high resolution map of the distribution of numerous unique markers along each of the organisms chromosomes may be obtained.

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As described in Example 56 below extended cDNAs (or genomic DNAs obtainable therefrom) may also be used to identify genes associated with a particular phenotype, such as hereditary disease or drug response.

3. Use of 5'ESTs or Sequences Obtained Therefrom or Fragments Thereof in Gene Identification

EXAMPLE 56

Identification of genes associated with hereditary diseases or drug response

This example illustrates an approach useful for the association of 5'ESTs (or cDNA or genomic DNA obtainable therefrom) with particular phenotypic characteristics. In this example, a particular 5'EST (or cDNA or genomic DNA obtainable therefrom) is used as a test probe to associate that 5'EST (or cDNA or genomic DNA obtainable therefrom) with a particular phenotypic characteristic.

5'ESTs (or cDNA or genomic DNA obtainable therefrom) are mapped to a particular location on a human chromosome using techniques such as those described in Examples 52 and 53 or other techniques known in the art. A search of Mendelian Inheritance in Man (McKusick in *Mendelian Inheritance in Man* (available on line through Johns Hopkins University Welch Medical Library) reveals the region of the human chromosome which contains the 5'EST (or cDNA or genomic DNA obtainable therefrom) to be a very gene rich region containing several known genes and several diseases or phenotypes for which genes have not been identified. The gene corresponding to this 5'EST (or cDNA or genomic DNA obtainable therefrom) thus becomes an immediate candidate for each of these genetic diseases.

Cells from patients with these diseases or phenotypes are isolated and expanded in culture. PCR primers from the 5'EST (or cDNA or genomic DNA obtainable therefrom) are used to screen genomic DNA, mRNA or cDNA obtained from the

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patients. 5'ESTs (or cDNA or genomic DNA obtainable therefrom) that are not amplified in the patients can be positively associated with a particular disease by further analysis. Alternatively, the PCR analysis may yield fragments of different lengths when the samples are derived from an individual having the phenotype associated with the disease than when the sample is derived from a healthy individual, indicating that the gene containing the 5'EST may be responsible for the genetic disease.

VI. Use of 5'EST (or cDNA or Genomic DNA Obtainable Therefrom) to Construct Vectors

The present 5'ESTs (or cDNA or genomic DNA obtainable therefrom) may also be used to construct secretion vectors capable of directing the secretion of the proteins encoded by genes therein. Such secretion vectors may facilitate the purification or enrichment of the proteins encoded by genes inserted therein by reducing the number of background proteins from which the desired protein must be purified or enriched.

15 Exemplary secretion vectors are described in Example 57 below.

1. Construction of Secretion Vectors

EXAMPLE 57

Construction of Secretion Vectors

The secretion vectors include a promoter capable of directing gene expression in the host cell, tissue, or organism of interest. Such promoters include the Rous Sarcoma Virus promoter, the SV40 promoter, the human cytomegalovirus promoter, and other promoters familiar to those skilled in the art.

A signal sequence from a 5' EST (or cDNAs or genomic DNAs obtainable therefrom) is operably linked to the promoter such that the mRNA transcribed from the promoter will direct the translation of the signal peptide. The host cell, tissue, or organism may be any cell, tissue, or organism which recognizes the signal peptide encoded by the signal sequence in the 5' EST (or cDNA or genomic DNA obtainable therefrom). Suitable hosts include mammalian cells, tissues or organisms, avian cells, tissues, or organisms, insect cells, tissues or organisms, or yeast.

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In addition, the secretion vector contains cloning sites for inserting genes encoding the proteins which are to be secreted. The cloning sites facilitate the cloning of the insert gene in frame with the signal sequence such that a fusion protein in which the signal peptide is fused to the protein encoded by the inserted gene is expressed from the mRNA transcribed from the promoter. The signal peptide directs the extracellular secretion of the fusion protein.

The secretion vector may be DNA or RNA and may integrate into the chromosome of the host, be stably maintained as an extrachromosomal replicon in the host, be an artificial chromosome, or be transiently present in the host. Many nucleic acid backbones suitable for use as secretion vectors are known to those skilled in the art, including retroviral vectors, SV40 vectors, Bovine Papilloma Virus vectors, yeast integrating plasmids, yeast episomal plasmids, yeast artificial chromosomes, human artificial chromosomes, P element vectors, baculovirus vectors, or bacterial plasmids capable of being transiently introduced into the host.

The secretion vector may also contain a polyA signal such that the polyA signal is located downstream of the gene inserted into the secretion vector.

After the gene encoding the protein for which secretion is desired is inserted into the secretion vector, the secretion vector is introduced into the host cell, tissue, or organism using calcium phosphate precipitation, DEAE-Dextran, electroporation, liposome-mediated transfection, viral particles or as naked DNA. The protein encoded by the inserted gene is then purified or enriched from the supernatant using conventional techniques such as ammonium sulfate precipitation, immunoprecipitation, immunochromatography, size exclusion chromatography, ion exchange chromatography, and HPLC. Alternatively, the secreted protein may be in a sufficiently enriched or pure state in the supernatant or growth media of the host to permit it to be used for its intended purpose without further enrichment.

The signal sequences may also be inserted into vectors designed for gene therapy. In such vectors, the signal sequence is operably linked to a promoter such that mRNA transcribed from the promoter encodes the signal peptide. A cloning site is located downstream of the signal sequence such that a gene encoding a protein whose secretion is desired may readily be inserted into the vector and fused to the signal sequence. The vector is introduced into an appropriate host cell. The protein expressed from the promoter is secreted extracellularly, thereby producing a therapeutic effect.

WO 99/06550 PCT/IB98/01232

102

The 5' ESTs may also be used to clone sequences located upstream of the 5' ESTs which are capable of regulating gene expression, including promoter sequences, enhancer sequences, and other upstream sequences which influence transcription or translation levels. Once identified and cloned, these upstream regulatory sequences may be used in expression vectors designed to direct the expression of an inserted gene in a desired spatial, temporal, developmental, or quantitative fashion. Example 58 describes a method for cloning sequences upstream of the extended cDNAs or 5' ESTs.

10 2. Identification of Upstream Sequences With Promoting or Regulatory Activities EXAMPLE 58

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Use of Extended cDNAs or 5' ESTs to Clone Upstream Sequences from Genomic DNA

Sequences derived from extended cDNAs or 5' ESTs may be used to isolate the promoters of the corresponding genes using chromosome walking techniques. In one chromosome walking technique, which utilizes the GenomeWalkerTM kit available from Clontech, five complete genomic DNA samples are each digested with a different restriction enzyme which has a 6 base recognition site and leaves a blunt end. Following digestion, oligonucleotide adapters are ligated to each end of the resulting genomic DNA fragments.

For each of the five genomic DNA libraries, a first PCR reaction is performed according to the manufacturer's instructions (which are incorporated herein by reference) using an outer adaptor primer provided in the kit and an outer gene specific primer. The gene specific primer should be selected to be specific for the extended cDNA or 5' EST of interest and should have a melting temperature, length, and location in the extended cDNA or 5'EST which is consistent with its use in PCR reactions. Each first PCR reaction contains 5 ng of genomic DNA, 5 µl of 10X Tth reaction buffer, 0.2 mM of each dNTP, 0.2 µM each of outer adaptor primer and outer gene specific primer, 1.1 mM of Mg(OAc)₂, and 1 µl of the Tth polymerase 50X mix in a total volume of 50 µl. The reaction cycle for the first PCR reaction is as follows: 1 min - 94°C / 2 sec - 94°C, 3 min - 72°C (7 cycles) / 2 sec - 94°C, 3 min - 67°C.

The product of the first PCR reaction is diluted and used as a template for a second PCR reaction according to the manufacturer's instructions using a pair of nested

WO 99/06550 PCT/IB98/01232

primers which are located internally on the amplicon resulting from the first PCR reaction. For example, 5 μl of the reaction product of the first PCR reaction mixture may be diluted 180 times. Reactions are made in a 50 μl volume having a composition identical to that of the first PCR reaction except the nested primers are used. The first nested primer is specific for the adaptor, and is provided with the GenomeWalker™ kit. The second nested primer is specific for the particular extended cDNA or 5' EST for which the promoter is to be cloned and should have a melting temperature, length, and location in the extended cDNA or 5' EST which is consistent with its use in PCR reactions. The reaction parameters of the second PCR reaction are as follows: 1 min - 94°C / 2 sec - 94°C, 3 min - 72°C (6 cycles) / 2 sec - 94°C, 3 min - 67°C (25 cycles) / 5 min - 67°C. The product of the second PCR reaction is purified, cloned, and sequenced using standard techniques.

Alternatively, two or more human genomic DNA libraries can be constructed by using two or more restriction enzymes. The digested genomic DNA is cloned into vectors which can be converted into single stranded, circular, or linear DNA. A biotinylated oligonucleotide comprising at least 15 nucleotides from the extended cDNA or 5' EST sequence is hybridized to the single stranded DNA. Hybrids between the biotinylated oligonucleotide and the single stranded DNA containing the extended cDNA or EST sequence are isolated as described in Example 29 above. Thereafter, the single stranded DNA containing the extended cDNA or EST sequence is released from the beads and converted into double stranded DNA using a primer specific for the extended cDNA or 5' EST sequence or a primer corresponding to a sequence included in the cloning vector. The resulting double stranded DNA is transformed into bacteria. DNAs containing the 5' EST or extended cDNA sequences are identified by colony PCR or colony hybridization.

Once the upstream genomic sequences have been cloned and sequenced as described above, prospective promoters and transcription start sites within the upstream sequences may be identified by comparing the sequences upstream of the extended cDNAs or 5' ESTs with databases containing known transcription start sites, transcription factor binding sites, or promoter sequences.

In addition, promoters in the upstream sequences may be identified using promoter reporter vectors as described in Example.

EXAMPLE 59

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Identification of Promoters in Cloned Upstream Sequences

The genomic sequences upstream of the extended cDNAs or 5' ESTs are cloned into a suitable promoter reporter vector, such as the pSEAP-Basic, pSEAP-Enhancer, pβgal-Basic, pβgal-Enhancer, or pEGFP-1 Promoter Reporter vectors available from Clontech. Briefly, each of these promoter reporter vectors include multiple cloning sites positioned upstream of a reporter gene encoding a readily assayable protein such as secreted alkaline phosphatase, β galactosidase, or green fluorescent protein. The sequences upstream of the extended cDNAs or 5' ESTs are inserted into the cloning sites upstream of the reporter gene in both orientations and introduced into an appropriate host cell. The level of reporter protein is assayed and compared to the level obtained from a vector which lacks an insert in the cloning site. The presence of an elevated expression level in the vector containing the insert with respect to the control vector indicates the presence of a promoter in the insert. If necessary, the upstream sequences can be cloned into vectors which contain an enhancer for augmenting transcription levels from weak promoter sequences. A significant level of expression above that observed with the vector lacking an insert indicates that a promoter sequence is present in the inserted upstream sequence.

Appropriate host cells for the promoter reporter vectors may be chosen based on the results of the above described determination of expression patterns of the extended cDNAs and ESTs. For example, if the expression pattern analysis indicates that the mRNA corresponding to a particular extended cDNA or 5' EST is expressed in fibroblasts, the promoter reporter vector may be introduced into a human fibroblast cell line.

Promoter sequences within the upstream genomic DNA may be further defined by constructing nested deletions in the upstream DNA using conventional techniques such as Exonuclease III digestion. The resulting deletion fragments can be inserted into the promoter reporter vector to determine whether the deletion has reduced or obliterated promoter activity. In this way, the boundaries of the promoters may be defined. If desired, potential individual regulatory sites within the promoter may be identified using site directed

mutagenesis or linker scanning to obliterate potential transcription factor binding sites within the promoter individually or in combination. The effects of these mutations on transcription levels may be determined by inserting the mutations into the cloning sites in the promoter reporter vectors.

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EXAMPLE 60

Cloning and Identification of Promoters

Using the method described in Example 58 above with 5' ESTs, sequences upstream of several genes were obtained. Using the primer pairs GGG AAG ATG GAG ATA GTA TTG CCT G (SEQ ID NO:29) and CTG CCA TGT ACA TGA TAG AGA GAT TC (SEQ ID NO:30), the promoter having the internal designation P13H2 (SEQ ID NO:31) was obtained.

Using the primer pairs GTA CCA GGGG ACT GTG ACC ATT GC (SEQ ID NO:32) and CTG TGA CCA TTG CTC CCA AGA GAG (SEQ ID NO:33), the promoter having the internal designation P15B4 (SEQ ID NO:34) was obtained.

Using the primer pairs CTG GGA TGG AAG GCA CGG TA (SEQ ID NO:35) and GAG ACC ACA CAG CTA GAC AA (SEQ ID NO:36), the promoter having the internal designation P29B6 (SEQ ID NO:37) was obtained.

Figure 4 provides a schematic description of the promoters isolated and the way they are assembled with the corresponding 5' tags. The upstream sequences were screened for the presence of motifs resembling transcription factor binding sites or known transcription start sites using the computer program MatInspector release 2.0, August 1996.

Table VII describes the transcription factor binding sites present in each of these promoters. The columns labeled matrice provides the name of the MatInspector matrix used. The column labeled position provides the 5' position of the promoter site. Numeration of the sequence starts from the transcription site as determined by matching the genomic sequence with the 5' EST sequence. The column labeled "orientation" indicates the DNA strand on which the site is found, with the + strand being the coding strand as determined by matching the genomic sequence with the sequence of the 5' EST. The column labeled "score" provides the MatInspector score found for this site. The column labeled "length" provides the length

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of the site in nucleotides. The column labeled "sequence" provides the sequence of the site found.

Bacterial clones containing plasmids containing the promoter sequences described above described above are presently stored in the inventor's laboratories under the internal identification numbers provided above. The inserts may be recovered from the deposited materials by growing an aliquot of the appropriate bacterial clone in the appropriate medium. The plasmid DNA can then be isolated using plasmid isolation procedures familiar to those skilled in the art such as alkaline lysis minipreps or large scale alkaline lysis plasmid isolation procedures. If desired the plasmid DNA may be further enriched by centrifugation on a cesium chloride gradient, size exclusion chromatography, or anion exchange chromatography. The plasmid DNA obtained using these procedures may then be manipulated using standard cloning techniques familiar to those skilled in the art. Alternatively, a PCR can be done with primers designed at both ends of the EST insertion. The PCR product which corresponds to the 5' EST can then be manipulated using standard cloning techniques familiar to those skilled in the art.

The promoters and other regulatory sequences located upstream of the extended cDNAs or 5' ESTs may be used to design expression vectors capable of directing the expression of an inserted gene in a desired spatial, temporal, developmental, or quantitative manner. A promoter capable of directing the desired spatial, temporal, developmental, and quantitative patterns may be selected using the results of the expression analysis described in Example 26 above. For example, if a promoter which confers a high level of expression in muscle is desired, the promoter sequence upstream of an extended cDNA or 5' EST derived from an mRNA which is expressed at a high level in muscle, as determined by the method of Example 26, may be used in the expression vector.

Preferably, the desired promoter is placed near multiple restriction sites to facilitate the cloning of the desired insert downstream of the promoter, such that the promoter is able to drive expression of the inserted gene. The promoter may be inserted in conventional nucleic acid backbones designed for extrachromosomal replication, integration into the host chromosomes or transient expression. Suitable backbones for the present expression vectors include retroviral backbones, backbones from eukaryotic episomes such as SV40 or Bovine Papilloma Virus, backbones from bacterial episomes, or artificial chromosomes.

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Preferably, the expression vectors also include a polyA signal downstream of the multiple restriction sites for directing the polyadenylation of mRNA transcribed from the gene inserted into the expression vector.

Following the identification of promoter sequences using the procedures of Examples 58-60, proteins which interact with the promoter may be identified as described in Example 61 below.

EXAMPLE 61

Identification of Proteins Which Interact with Promoter Sequences, Upstream Regulatory Sequences, or mRNA

Sequences within the promoter region which are likely to bind transcription factors may be identified by homology to known transcription factor binding sites or through conventional mutagenesis or deletion analyses of reporter plasmids containing the promoter sequence. For example, deletions may be made in a reporter plasmid containing the promoter sequence of interest operably linked to an assayable reporter gene. The reporter plasmids carrying various deletions within the promoter region are transfected into an appropriate host cell and the effects of the deletions on expression levels is assessed. Transcription factor binding sites within the regions in which deletions reduce expression levels may be further localized using site directed mutagenesis, linker scanning analysis, or other techniques familiar to those skilled in the art.

Nucleic acids encoding proteins which interact with sequences in the promoter may be identified using one-hybrid systems such as those described in the manual accompanying the Matchmaker One-Hybrid System kit available from Clontech (Catalog No. K1603-1), the disclosure of which is incorporated herein by reference. Briefly, the Matchmaker One-hybrid system is used as follows. The target sequence for which it is desired to identify binding proteins is cloned upstream of a selectable reporter gene and integrated into the yeast genome. Preferably, multiple copies of the target sequences are inserted into the reporter plasmid in tandem. A library comprised of fusions between cDNAs to be evaluated for the ability to bind to the promoter and the activation domain of a yeast transcription factor, such as GAL4, is transformed into the yeast strain containing the integrated reporter sequence. The yeast are plated on selective media to

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select cells expressing the selectable marker linked to the promoter sequence. The colonies which grow on the selective media contain genes encoding proteins which bind the target sequence. The inserts in the genes encoding the fusion proteins are further characterized by sequencing. In addition, the inserts may be inserted into expression vectors or *in vitro* transcription vectors. Binding of the polypeptides encoded by the inserts to the promoter DNA may be confirmed by techniques familiar to those skilled in the art, such as gel shift analysis or DNAse protection analysis.

VII. Use of 5' ESTs (or cDNAs or Genomic DNAs Obtainable Therefrom) in Gene Therapy

The present invention also comprises the use of 5'ESTs (or cDNA or genomic DNA obtainable therefrom) in gene therapy strategies, including antisense and triple helix strategies as described in Examples 62 and 63 below. In antisense approaches, nucleic acid sequences complementary to an mRNA are hybridized to the mRNA intracellularly, thereby blocking the expression of the protein encoded by the mRNA. The antisense sequences may prevent gene expression through a variety of mechanisms. For example, the antisense sequences may inhibit the ability of ribosomes to translate the mRNA. Alternatively, the antisense sequences may block transport of the mRNA from the nucleus to the cytoplasm, thereby limiting the amount of mRNA available for translation. Another mechanism through which antisense sequences may inhibit gene expression is by interfering with mRNA splicing. In yet another strategy, the antisense nucleic acid may be incorporated in a ribozyme capable of specifically cleaving the target mRNA.

EXAMPLE 62

Preparation and Use of Antisense Oligonucleotides

The antisense nucleic acid molecules to be used in gene therapy may be either DNA or RNA sequences. They may comprise a sequence complementary to the sequence of the 5'EST (or cDNA or genomic DNA obtainable therefrom). The antisense nucleic acids should have a length and melting temperature sufficient to permit formation of an intracellular duplex with sufficient stability to inhibit the expression of the mRNA in the duplex. Strategies for designing antisense nucleic acids suitable for use in gene therapy are disclosed in Green et

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al., Ann. Rev. Biochem. 55:569-597, 1986; and Izant and Weintraub, Cell 36:1007-1015, 1984, which are hereby incorporated by reference.

In some strategies, antisense molecules are obtained from a nucleotide sequence encoding a protein by reversing the orientation of the coding region with respect to a promoter so as to transcribe the opposite strand from that which is normally transcribed in the cell. The antisense molecules may be transcribed using *in vitro* transcription systems such as those which employ T7 or SP6 polymerase to generate the transcript. Another approach involves transcription of the antisense nucleic acids *in vivo* by operably linking DNA containing the antisense sequence to a promoter in an expression vector.

Alternatively, oligonucleotides which are complementary to the strand normally transcribed in the cell may be synthesized *in vitro*. Thus, the antisense nucleic acids are complementary to the corresponding mRNA and are capable of hybridizing to the mRNA to create a duplex. In some embodiments, the antisense sequences may contain modified sugar phosphate backbones to increase stability and make them less sensitive to RNase activity. Examples of modifications suitable for use in antisense strategies are described by Rossi *et al.*, *Pharmacol. Ther.* 50(2):245-254, 1991, which is hereby incorporated by reference.

Various types of antisense oligonucleotides complementary to the sequence of the 5'EST (or cDNA or genomic DNA obtainable therefrom) may be used. In one preferred embodiment, stable and semi-stable antisense oligonucleotides described in International Application No. PCT WO94/23026, hereby incorporated by reference, are used. In these molecules, the 3' end or both the 3' and 5' ends are engaged in intramolecular hydrogen bonding between complementary base pairs. These molecules are better able to withstand exonuclease attacks and exhibit increased stability compared to conventional antisense oligonucleotides.

In another preferred embodiment, the antisense oligodeoxynucleotides against herpes simplex virus types 1 and 2 described in International Application No. WO 95/04141, hereby incorporated by reference, are used.

In yet another preferred embodiment, the covalently cross-linked antisense oligonucleotides described in International Application No. WO 96/31523, hereby incorporated by reference, are used. These double- or single-stranded oligonucleotides comprise one or more, respectively, inter- or intra-oligonucleotide covalent cross-linkages,

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wherein the linkage consists of an amide bond between a primary amine group of one strand and a carboxyl group of the other strand or of the same strand, respectively, the primary amine group being directly substituted in the 2' position of the strand nucleotide monosaccharide ring, and the carboxyl group being carried by an aliphatic spacer group substituted on a nucleotide or nucleotide analog of the other strand or the same strand, respectively.

The antisense oligodeoxynucleotides and oligonucleotides disclosed in International Application No. WO 92/18522, incorporated by reference, may also be used. These molecules are stable to degradation and contain at least one transcription control recognition sequence which binds to control proteins and are effective as decoys therefore. These molecules may contain "hairpin" structures, "dumbbell" structures, "modified dumbbell" structures, "cross-linked" decoy structures and "loop" structures.

In another preferred embodiment, the cyclic double-stranded oligonucleotides described in European Patent Application No. 0 572 287 A2, hereby incorporated by reference are used. These ligated oligonucleotide "dumbbells" contain the binding site for a transcription factor and inhibit expression of the gene under control of the transcription factor by sequestering the factor.

Use of the closed antisense oligonucleotides disclosed in International Application No. WO 92/19732, hereby incorporated by reference, is also contemplated. Because these molecules have no free ends, they are more resistant to degradation by exonucleases than are conventional oligonucleotides. These oligonucleotides may be multifunctional, interacting with several regions which are not adjacent to the target mRNA.

The appropriate level of antisense nucleic acids required to inhibit gene expression may be determined using *in vitro* expression analysis. The antisense molecule may be introduced into the cells by diffusion, injection, infection, transfection or h-region-mediated import using procedures known in the art. For example, the antisense nucleic acids can be introduced into the body as a bare or naked oligonucleotide, oligonucleotide encapsulated in lipid, oligonucleotide sequence encapsidated by viral protein, or as an oligonucleotide operably linked to a promoter contained in an expression vector. The expression vector may be any of a variety of expression vectors known in the art, including retroviral or viral vectors,

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vectors capable of extrachromosomal replication, or integrating vectors. The vectors may be DNA or RNA.

The antisense molecules are introduced onto cell samples at a number of different concentrations preferably between $1\times10^{-10}M$ to $1\times10^{-4}M$. Once the minimum concentration that can adequately control gene expression is identified, the optimized dose is translated into a dosage suitable for use *in vivo*. For example, an inhibiting concentration in culture of 1×10^{-7} translates into a dose of approximately 0.6 mg/kg bodyweight. Levels of oligonucleotide approaching 100 mg/kg bodyweight or higher may be possible after testing the toxicity of the oligonucleotide in laboratory animals. It is additionally contemplated that cells from the vertebrate are removed, treated with the antisense oligonucleotide, and reintroduced into the vertebrate.

It is further contemplated that the antisense oligonucleotide sequence is incorporated into a ribozyme sequence to enable the antisense to specifically bind and cleave its target mRNA. For technical applications of ribozyme and antisense oligonucleotides see Rossi et al., supra.

In a preferred application of this invention, the polypeptide encoded by the gene is first identified, so that the effectiveness of antisense inhibition on translation can be monitored using techniques that include but are not limited to antibody-mediated tests such as RIAs and ELISA, functional assays, or radiolabeling.

The 5' ESTs of the present invention (or cDNAs or genomic DNAs obtainable therefrom) may also be used in gene therapy approaches based on intracellular triple helix formation. Triple helix oligonucleotides are used to inhibit transcription from a genome. They are particularly useful for studying alterations in cell activity as it is associated with a particular gene. The 5' EST sequences (or cDNAs or genomic DNAs obtainable therefrom) of the present invention or, more preferably, a portion of those sequences, can be used to inhibit gene expression in individuals having diseases associated with expression of a particular gene. Similarly, a portion of 5' EST sequences (or cDNAs or genomic DNAs obtainable therefrom) can be used to study the effect of inhibiting transcription of a particular gene within a cell. Traditionally, homopurine sequences were considered the most useful for triple helix strategies. However, homopyrimidine sequences can also inhibit gene expression. Such homopyrimidine oligonucleotides bind to the major groove

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homopurine:homopyrimidine sequences. Thus, both types of sequences from the 5'EST or from the gene corresponding to the 5'EST are contemplated within the scope of this invention.

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EXAMPLE 63

Preparation and Use of Triple Helix Probes

The sequences of the 5' ESTs (or cDNAs or genomic DNAs obtainable therefrom) are scanned to identify 10-mer to 20-mer homopyrimidine or homopurine stretches which could be used in triple-helix based strategies for inhibiting gene expression. Following identification of candidate homopyrimidine or homopurine stretches, their efficiency in inhibiting gene expression is assessed by introducing varying amounts of oligonucleotides containing the candidate sequences into tissue culture cells which normally express the target gene. The oligonucleotides may be prepared on an oligonucleotide synthesizer or they may be purchased commercially from a company specializing in custom oligonucleotide synthesis, such as GENSET, Paris, France.

The oligonucleotides may be introduced into the cells using a variety of methods known to those skilled in the art, including but not limited to calcium phosphate precipitation, DEAE-Dextran, electroporation, liposome-mediated transfection or native uptake.

Treated cells are monitored for altered cell function or reduced gene expression using techniques such as Northern blotting, RNase protection assays, or PCR based strategies to monitor the transcription levels of the target gene in cells which have been treated with the oligonucleotide. The cell functions to be monitored are predicted based upon the homologies of the target gene corresponding to the extended cDNA from which the oligonucleotide was derived with known gene sequences that have been associated with a particular function. The cell functions can also be predicted based on the presence of abnormal physiologies within cells derived from individuals with a particular inherited disease, particularly when the extended cDNA is associated with the disease using techniques described in Example 56.

The oligonucleotides which are effective in inhibiting gene expression in tissue culture cells may then be introduced *in vivo* using the techniques described above and in Example 62 at a dosage calculated based on the *in vitro* results, as described in Example 62.

113

In some embodiments, the natural (beta) anomers of the oligonucleotide units can be replaced with alpha anomers to render the oligonucleotide more resistant to nucleases. Further, an intercalating agent such as ethidium bromide, or the like, can be attached to the 3' end of the alpha oligonucleotide to stabilize the triple helix. For information on the generation of oligonucleotides suitable for triple helix formation see Griffin *et al.*, *Science* 245:967-971, 1989, which is hereby incorporated by this reference.

EXAMPLE 64

Use of cDNAs Obtained Using the 5' ESTs to Express an Encoded Protein in a Host

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The cDNAs obtained as described above using the 5' ESTs of the present invention may also be used to express an encoded protein in a host organism to produce a beneficial effect. In such procedures, the encoded protein may be transiently expressed in the host organism or stably expressed in the host organism. The encoded protein may have any of the activities described above. The encoded protein may be a protein which the host organism lacks or, alternatively, the encoded protein may augment the existing levels of the protein in the host organism.

A full length extended cDNA encoding the signal peptide and the mature protein, or an extended cDNA encoding only the mature protein is introduced into the host organism. The extended cDNA may be introduced into the host organism using a variety of techniques known to those of skill in the art. For example, the extended cDNA may be injected into the host organism as naked DNA such that the encoded protein is expressed in the host organism, thereby producing a beneficial effect.

Alternatively, the extended cDNA may be cloned into an expression vector downstream of a promoter which is active in the host organism. The expression vector may be any of the expression vectors designed for use in gene therapy, including viral or retroviral vectors. The expression vector may be directly introduced into the host organism such that the encoded protein is expressed in the host organism to produce a beneficial effect. In another approach, the expression vector may be introduced into cells *in vitro*. Cells containing the expression vector are thereafter selected and introduced into the host organism, where they express the encoded protein to produce a beneficial effect.

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EXAMPLE 65

<u>Use of Signal Peptides Encoded by 5' ESTs or Sequences obtained Therefrom</u> <u>to Import Proteins Into Cells</u>

The short core hydrophobic region (h) of signal peptides encoded by the 5'ESTS or extended cDNAs derived from SEQ ID NOs: 38-315 may also be used as a carrier to import a peptide or a protein of interest, so-called cargo, into tissue culture cells (Lin et al., J. Biol. Chem., 270: 14225-14258, 1995; Du et al., J. Peptide Res., 51: 235-243, 1998; Rojas et al., Nature Biotech., 16: 370-375, 1998).

When cell permeable peptides of limited size (approximately up to 25 amino acids) are to be translocated across cell membrane, chemical synthesis may be used in order to add the h region to either the C-terminus or the N-terminus to the cargo peptide of interest. Alternatively, when longer peptides or proteins are to be imported into cells, nucleic acids can be genetically engineered, using techniques familiar to those skilled in the art, in order to link the extended cDNA sequence encoding the h region to the 5' or the 3' end of a DNA sequence coding for a cargo polypeptide. Such genetically engineered nucleic acids are then translated either *in vitro* or *in vivo* after transfection into appropriate cells, using conventional techniques to produce the resulting cell permeable polypeptide. Suitable hosts cells are then simply incubated with the cell permeable polypeptide which is then translocated across the membrane.

This method may be applied to study diverse intracellular functions and cellular processes. For instance, it has been used to probe functionally relevant domains of intracellular proteins and to examine protein-protein interactions involved in signal transduction pathways (Lin et al., supra; Lin et al., J. Biol. Chem., 271: 5305-5308, 1996; Rojas et al., J. Biol. Chem., 271: 27456-27461, 1996; Liu et al., Proc. Natl. Acad. Sci. USA, 93: 11819-11824, 1996; Rojas et al., Bioch. Biophys. Res. Commun., 234: 675-680, 1997).

Such techniques may be used in cellular therapy to import proteins producing therapeutic effects. For instance, cells isolated from a patient may be treated with imported therapeutic proteins and then re-introduced into the host organism.

Alternatively, the h region of signal peptides of the present invention could be used in combination with a nuclear localization signal to deliver nucleic acids into cell nucleus. Such oligonucleotides may be antisense oligonucleotides or oligonucleotides designed to form

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triple helixes, as described in examples 62 and 63 respectively, in order to inhibit processing and/or maturation of a target cellular RNA.

As discussed above, the cDNAs or portions thereof obtained using the 5' ESTs of the present invention can be used for various purposes. The polynucleotides can be used to express recombinant protein for analysis, characterization or therapeutic use; as markers for tissues in which the corresponding protein is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in disease states); as molecular weight markers on Southern gels; as chromosome markers or tags (when labeled) to identify chromosomes or to map related gene positions; to compare with endogenous DNA sequences in patients to identify potential genetic disorders; as probes to hybridize and thus discover novel, related DNA sequences; as a source of information to derive PCR primers for genetic fingerprinting; for selecting and making oligomers for attachment to a "gene chip" or other support, including for examination for expression patterns; to raise anti-protein antibodies using DNA immunization techniques; and as an antigen to raise anti-DNA antibodies or elicit another immune response. Where the polynucleotide encodes a protein which binds or potentially binds to another protein (such as, for example, in a receptor-ligand interaction), the polynucleotide can also be used in interaction trap assays (such as, for example, that described in Gyuris et al., Cell 75:791-803, 1993, the disclosure of which is hereby incorporated by reference) to identify polynucleotides encoding the other protein with which binding occurs or to identify inhibitors of the binding interaction.

The proteins or polypeptides provided by the present invention can similarly be used in assays to determine biological activity, including in a panel of multiple proteins for high-throughput screening; to raise antibodies or to elicit another immune response; as a reagent (including the labeled reagent) in assays designed to quantitatively determine levels of the protein (or its receptor) in biological fluids; as markers for tissues in which the corresponding protein is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in a disease state); and, of course, to isolate correlative receptors or ligands. Where the protein binds or potentially binds to another protein (such as, for example, in a receptor-ligand interaction), the protein can be used to identify the other protein with which binding occurs or to identify inhibitors of the binding interaction. Proteins

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involved in these binding interactions can also be used to screen for peptide or small molecule inhibitors or agonists of the binding interaction.

Any or all of these research utilities are capable of being developed into reagent grade or kit format for commercialization as research products.

Methods for performing the uses listed above are well known to those skilled in the art. References disclosing such methods include without limitation *Molecular Cloning*; A Laboratory Manual, 2d ed., Cold Spring Harbor Laboratory Press, Sambrook, Fritsch and Maniatis eds., 1989, and Methods in Enzymology; Guide to Molecular Cloning Techniques, Academic Press, Berger and Kimmel eds., 1987.

Polynucleotides and proteins of the present invention can also be used as nutritional sources or supplements. Such uses include without limitation use as a protein or amino acid supplement, use as a carbon source, use as a nitrogen source and use as a source of carbohydrate. In such cases the protein or polynucleotide of the invention can be added to the feed of a particular organism or can be administered as a separate solid or liquid preparation, such as in the form of powder, pills, solutions, suspensions or capsules. In the case of microorganisms, the protein or polynucleotide of the invention can be added to the medium in or on which the microorganism is cultured.

Although this invention has been described in terms of certain preferred embodiments, other embodiments which will be apparent to those of ordinary skill in the art in view of the disclosure herein are also within the scope of this invention. Accordingly, the scope of the invention is intended to be defined only by reference to the appended claims. All documents cited herein are incorporated herein by reference in their entirety.

	Search characteristic	cteristic	Selection	Selection Characteristics	,
Step	Program	Strand	Parameters	Identity (%)	Length (bp)
miscellanaeous	blastn	both	S=61 X=16	06	17
tRNA	fasta	both		80	90
rRNA	blastn	both	S=108	80	40
mtRNA	blastn	both	S=108	80	40
Procaryotic	blastn	both	S=144	06	40
Fungal	blastn	both	S=144	06	40
Alu	fasta*	both	•	70	40
	blastn	both	S=72	70	40
Repeats	blastn	both	S=72	70	40
Promoters	biastn	top	S=54 X=16	06	15†
Vertebrate	fasta*	both	S=108	06	30
ESTs	blastn	both	S=108 X=16	06	30
Proteins	blastx¤	top	E = 0.001	•	-

Table 1: Parameters used for each step of EST analysis

use "Quick Fast" Database scanner
 alignement further constrained to begin closer than 10bp to EST\5' end
 using BLOSUM62 substitution matrix

TABLE II

		IAI	DLE II	
SEQ. ID		VON HEIJNE	TISSUE	
<u>NO.</u>	CATEGORY	SCORE		INTERNAL
		<u> </u>	SOURCE	DESIGNATION
ID38	new	11.4	Canage	
ID39	new	11.3	Cancerous prostate	76-36-2-G4-PU
ID40	new	11.5	Normal prostate	78-26-1-A7-PU
ID41	new	10.7	Normal prostate	78-4-3-G8-PU
ID42	new	10.7	Hypertrophic prostate	77-16-3-D7 - PU
ID43	new	10.6	Hypertrophic prostate	77-7-1-H9-PU
ID44	new	10.6	Hypertrophic prostate	77-42-1-D10-PU
ID45	new	10.4	Cancerous prostate	76-34-4-C6-PU
ID46	new	10.2	Normal prostate	78-31-3-B8-PU
ID47	new	10.2	Normal prostate	78-38-1-C10-PU
ID48	new	9	Cancerous prostate	76-16-4-D5-PU
ID49	new	8.8	Hypertrophic prostate	77-38-2-B9-PU
ID50	new	8.6	Normal prostate	78-30-1 -G 12-PU
ID51	new	8.5	Prostate	60-17-1-F1-PU
ID52	new	8.3	Prostate	60-17-3 -G 8-PU
ID53	new	8.3	Normal prostate	78-8-2-H8-PU
ID54	new	8.3 8.3	Normal prostate	78-26-2-A1-PU
ID55	new	8.2	Cancerous prostate	76-23-2-B10 - PU
ID56	new	8.1	Cancerous prostate	76-23-4-H9 -PU
ID57	new		Normal prostate	78-44-2-C3-PU
ID58	new	8 8	Hypertrophic prostate	77-37-1-H3-PU
ID59	new	7.8	Normal prostate	78-35-2-G12-PU
ID60	new		Normal prostate	78-17-4-G2-PU
ID61	new	7.7	Normal prostate	78-5-4-F7-PU
ID62	new	7.6 7.6	Normal prostate	78-16-3-E2-PU
ID63	new	-	Hypertrophic prostate	77-5-1-B6-PU
ID64	new	7.6 7.5	Normal prostate	78-26-1-B5-PU
ID65	new	7.5 7.5	Cancerous prostate	76-12-1-B1-PU
ID66	new	7.5 7.3	Normal prostate	78-4-4-E7-PU
ID67	new	7.2	Hypertrophic prostate	77-11-1-A3-PU
ID68	new	7.2	Hypertrophic prostate	77-5-4-G9-PU
ID69	new	7.2	Normal prostate	78-23-4-H11-PU
ID70 ·	new	7.2	Hypertrophic prostate	77-39-3-H7-PU
ID71	new	7.2	Cancerous prostate	76-23-4-H2-PU
ID72	new	7.2	Cancerous prostate	76-24-1-F8-PU
ID73	new	7	Normal prostate	78-39-4-D2-PU
ID74		7	Normal prostate	78-28-3-D2-PU
ID75	new	7	Normal prostate	78-29-3-H11-PU
ID76	new	7	Normal prostate	78-40-3-G2-PU
ID77	new	7	Cancerous prostate	76-1-2-F8-PU
ID78	new	7	Normal prostate	78-13-4-B10-PU
ID79	new	6.9	Cancerous prostate	76-12-1-A9-PU
ID80	new	6.9	Normal prostate	78-20-3-C11-PU
ID81	new	6.9	Cancerous prostate	76-9-2-D10-PU
ID81 ID82	new	6.8	Normal prostate	78-6-2-D12-PU
ID83	new	6.7	Hypertrophic prostate	77-10-1-C8-PU
ID83	new	6.7	Cancerous prostate	76-13-2-F11 -P U
ID85	new	6.7	Cancerous prostate	76-4-1-G5-PU
ID86	new	6.5	Normal prostate	78-3-4-B8-PU
ID87	new	6.4	Prostate	60-11-3-G2-PU
ID88	new	6.3	Normal prostate	78-25-1-G5-PU
1700	new	6.3	Normal prostate	78-2-2-G5-PU
				- -

CEO ID				
SEQ. ID	CATEGORY	VON HEIJNE	TISSUE	INTERNAL
<u>NO.</u>	CATEGORI	SCORE	SOURCE	<u>DESIGNATION</u>
ID89	new	6.3	Cancerous prostate	76 7 2 A 1 DII
ID90	new	6.3	Hypertrophic prostate	76-7-3-A1-PU
ID91	new	6.2	Normal prostate	77-5-1-C2-PU
ID92	new	6.1	Normal prostate	78-49-2-A11-PU
ID93	new	6	<u> </u>	78-7-1-B9-PU
ID94	new	6	Normal prostate	78-39-4-G3-PU
ID95	new	5.9	Normal prostate	78-32-2-H6-PU
ID96	new	5.9	Cancerous prostate	76-30-3-H2-PU
ID97	new	5.9	Normal prostate	78-24-3-H4-PU
ID97 ID98	new	5.8	Cancerous prostate	76-43-3-B6-PU
ID99	new	5.8	Prostate	60-16-3-A3-PU
ID100			Cancerous prostate	76-20-4-C11-PU
ID100	new	5.7	Cancerous prostate	76-11-1-C5-PU
	new	5.7	Hypertrophic prostate	77-37-3-C1-PU
ID102	new	5.7	Prostate	60-13-2-B5-PU
ID103	new	5.7	Normal prostate	78-49-4-E4-PU
ID104	new	5.6	Normal prostate	78-37-4-C11-PU
ID105	new	5.6	Prostate	60-17-1-D8-PU
ID106	new	5.5	Normal prostate	78-36-3-D7-PU
ID107	new	5.5	Cancerous prostate	76-24-3-E11-PU
ID108	new	5.5	Prostate	60-14-2-A7-PU
ID109	new	5.4	Hypertrophic prostate	77-10-4-F9-PU
ID110	new	5.3	Cancerous prostate	76-23-3-G5-PU
D111	new	5.3	Normal prostate	78-42-3-D3-PU
ID112	new	5.3	Prostate	60-12-1-H1-PU
ID113	new	5.3	Hypertrophic prostate	77-5-2-A3-PU
ID114	new	5.2	Normal prostate	78-37-2-G12-PU
ID115	new	5.2	Cancerous prostate	76-39-2-H1-PU
ID116	new	5.1	Prostate	60-12-3-C2-PU
ID117	new	5.1	Normal prostate	78-25-1-F11-PU
ID118	new	5.1	Normal prostate	78-36-2-C10-PU
ID119	new	5.1	Hypertrophic prostate	77-13-1-B7-PU
ID120	new	5.1	Hypertrophic prostate	77-4-4-H7-PU
ID121	new	5	Normal prostate	78-33-4-F9-PU
ID122	new	5	Cancerous prostate	76-21-1-D5-PU
ID123	new	4.8	Normal prostate	78-3-4-B3-PU
ID124	new	4.8	Cancerous prostate	76-29-4-B3-PU
ID125	new	4.8	Normal prostate	78-46-3-C6-PU
ID126	new	4.8	Hypertrophic prostate	77-13-3-F8-PU
ID127	new	4.7	Cancerous prostate	76-12-4-C3 - PU
ID128	new	4.7	Cancerous prostate	76-34-4-C1-PU
ID129	new	4.7	Normal prostate	78-42-4-D2-PU
ID130	new	4.7	Cancerous prostate	76-38-2-H9-PU
ID131	new	4.6	Normal prostate	78-49-4-B5-PU
ID132	new	4.6	Cancerous prostate	76-1-1-E3-PU
ID133	new	4.6	Normal prostate	78-46-3-C4-PU
ID134	new	4.5	Cancerous prostate	76-22-2-D2-PU
ID135	new	4.5	Prostate	
ID136	new	4.5	Normal prostate	60-11-4-F6-PU
ID137	new	4.4	Prostate	78-32-2-G1-PU
ID138	new	4.4		60-14-3-C7-PU
ID138	new	4.4	Hypertrophic prostate Normal prostate	77-3-4-H3-PU
ID140	new	4.3		78-36-4-E12-PU
ID140 ID141	new	4.3	Hypertrophic prostate	77-12-1-A9-PU
	110-17	7.3	Normal prostate	78-23-2-H3-PU

SEQ. ID		MONTHS -		
NO.	CATEGORY	VON HEIJNE	TISSUE	INTERNAL
	CHILOOKI	SCORE	SOURCE	DESIGNATION
ID142	new	4.2	Cancerous promotes	
ID143	new	4.2	Cancerous prostate Normal prostate	76-39-3-C11-PU
ID144	new	4.2	Cancerous prostate	78-23-3-D10-PU
ID145	new	4.2	Normal prostate	76-32-2-B7-PU
ID146	new	4.2	Prostate	78-40-1-G9-PU
ID147	new	4.1	Cancerous prostate	60-12-1-E11-PU 76-27-3-A6-PU
ID148	new	4	Cancerous prostate	76-43-3-B2-PU
ID149	new	4	Normal prostate	78-18-3-B4-PU
ID150	new	4	Normal prostate	78-41-2-D11-PU
ID151	new	4	Normal prostate	78-34-2-G9-PU
ID152	new	4	Normal prostate	78-4-3-G2-PU
ID153	new	4	Hypertrophic prostate	77-22-2-G2-PU
ID154	new	3.9	Cancerous prostate	76-4-4-F6-PU
ID155	new	3.9	Hypertrophic prostate	77-40-3-E10-PU
ID156	new	3.9	Normal prostate	78-10-1-H5-PU
ID157	new	3.9	Normal prostate	78-6-2-E3-PU
ID158	new	3.9	Hypertrophic prostate	77-20-3-E5-PU
ID159 ID160	new	3.9	Normal prostate	78-38-2-B5-PU
ID160 ID161	new	3.8	Prostate	60-11-2-G12-PU
ID161	new	3.8	Cancerous prostate	76-44-3-E8-PU
ID162 ID163	new	3.8	Normal prostate	78-41-3-A2-PU
ID163	new	3.7	Cancerous prostate	76-20-4-E7-PU
ID165	new	3.7	Cancerous prostate	76-17-1-E4-PU
ID166	new	3.7	Normal prostate	78-5-2-D2-PU
ID167	new	3.7	Prostate	60-11 - 3-B11-PU
ID168	new new	3.7	Hypertrophic prostate	77-21-2-F1-PU
ID169	new	3.6	Prostate	60-12-1-A5-PU
ID170	new	3.6	Cancerous prostate	76-18-2-G12-PU
ID171	new	3.6 3.6	Normal prostate	78-7-1-G5-PU
ID172	new	3.5	Cancerous prostate	76-37-4-A5-PU
ID173	new	3.5 3.5	Normal prostate	78-50-4-A2-PU
ID174	new	3.5	Normal prostate	78-43-2-H10-PU
ID175	new	3.5	Normal prostate	78-44-3-B6-PU
ID176	new	3.5	Cancerous prostate	76-10-1-D6-PU
ID177	new	3.5	Prostate	60-11-4-F2-PU
ID178	ext-est-not-vrt	14.8	Cancerous prostate Normal prostate	76-45-2-B12-PU
ID179	ext-est-not-vrt	13.6	Normal prostate	78-34-3-D9-PU
ID180	ext-est-not-vrt	12.7	Normal prostate	78-46-4-F4-PU
ID181	ext-est-not-vrt	8.8	Prostate	78-8-3-D9-PU
ID182	ext-est-not-vrt	8.5	Normal prostate	60-15-4-F6-PU
ID183	ext-est-not-vrt	7.3	Normal prostate	78-8-3-E6-PU
ID184	ext-est-not-vrt	7.1	Cancerous prostate	78-7-3-A4-PU
ID185	ext-est-not-vrt	6.6	Cancerous prostate	76-33-2-F5-PU
ID186	ext-est-not-vrt	6.3	Normal prostate	76-34-4-G12-PU
ID187	ext-est-not-vrt	5.9	Normal prostate	78-13-1-H7-PU
ID188	ext-est-not-vrt	5.9	Normal prostate	78-49-3-B11-PU
ID189	ext-est-not-vrt	5.5	Cancerous prostate	78-42-2-A10-PU 76-7-4-D9-PU
ID190	ext-est-not-vrt	5.2	Normal prostate	78-40-3-B12-PU
ID191	ext-est-not-vrt	5	Hypertrophic prostate	77-36-1-G2-PU
ID192	ext-est-not-vrt	4.8	Prostate	60-17-3-H11-PU
ID193	ext-est-not-vrt	4.4	Normal prostate	78-28-3-E4-PU
ID194	ext-est-not-vrt	4.1	Cancerous prostate	76-28-2-H5-PU
			-	. 5 20 2-1D-10

SEQ. ID		VON HEIJNE	TISSUE	INTERNAL
<u>NO.</u>	<u>CATEGORY</u>	_SCORE	SOURCE	DESIGNATION
ID195	ext-est-not-vrt	4.1	Normal prostate	78-27-1-D11-PU
ID196	ext-est-not-vrt	3.9	Cancerous prostate	76-42-2-B5-PU
ID 197	ext-est-not-vrt	3.9	Hypertrophic prostate	77-39-3-F8-PU
ID198	ext-est-not-vrt	3.7	Cancerous prostate	76-43-1-G9-PU
ID 199	est-not-ext	13.8	Normal prostate	78-40-1-B10-PU
ID200	est-not-ext	13.4	Cancerous prostate	76-15-1-F4-PU
ID201	est-not-ext	13	Cancerous prostate	76-45-4-E7-PU
ID202	est-not-ext	11.6	Normal prostate	78-26-2-H7-PU
ID203	est-not-ext	11.2	Normal prostate	78-21-1-B7-PU
ID204	est-not-ext	11.2	Cancerous prostate	76-40-2-F5-PU
ID205	est-not-ext	10.6	Cancerous prostate	76-29-2-G8-PU
ID206	est-not-ext	10.5	Hypertrophic prostate	77-23-4-H11-PU
ID207	est-not-ext .	10.3	Normal prostate	78-48-1-F10-PU
ID208	est-not-ext	9.5	Cancerous prostate	76-41-4-G9-PU
ID209	est-not-ext	9.3	Hypertrophic prostate	77-3-3-C10-PU
ID210	est-not-ext	9.1	Cancerous prostate	76-45-4-C8-PU
ID211	est-not-ext	8.8	Normal prostate	78-50-4-C10-PU
ID212	est-not-ext	8.8	Normal prostate	78-38-4-F7-PU
ID213	est-not-ext	8.6	Cancerous prostate	
ID214	est-not-ext	8.6	Normal prostate	76-16-4-C9-PU
ID215	est-not-ext	8.4	Cancerous prostate	78-49-2-D10-PU
ID216	est-not-ext	7.9	Normal prostate	76-1-1-H7-PU
ID217	est-not-ext	7.9 7.9	•	78-4-2-F10-PU
ID218	est-not-ext	7.7 7.7	Normal prostate	78-46-3-B6-PU
ID219	est-not-ext	7.6 7.6	Normal prostate	78-7-1-F2-PU
ID220	est-not-ext	7.6 7.6	Normal prostate	78-35-2-D3-PU
ID221	est-not-ext	7.6	Cancerous prostate	76-20-2-G7-PU
ID222	est-not-ext	7.6 7.5	Normal prostate	78-39-1-E11-PU
ID222 ID223			Cancerous prostate	76-4-4-C2-PU
ID223 ID224	est-not-ext	7.1 7	Normal prostate	78-48-2-F6-PU
ID224 ID225	est-not-ext	-	Cancerous prostate	76-32-4-A10-PU
	est-not-ext	6.8	Cancerous prostate	76-39-1-E7-PU
ID226	est-not-ext	6.7	Cancerous prostate	76-29-4-E1-PU
ID227	est-not-ext	6.7	Normal prostate	78-28-4-B9-PU
ID228	est-not-ext	6.7	Normal prostate	78-37-4-B2-PU
ID229 T	est-not-ext	6.7	Normal prostate	78-50-2-E12 <i>-</i> PU
ID230	est-not-ext	6.7	Hypertrophic prostate	77-21-2-F8-PU
ID231	est-not-ext	6.6	Normal prostate	78-27-4-E2-PU
ID232	est-not-ext	6.5	Normal prostate	78-45-4-G12-PU
ID233	est-not-ext	6.3	Cancerous prostate	76-7-4-H8-PU
ID234	est-not-ext	6.3	Normal prostate	78-23-1-D10-PU
ID235	est-not-ext	6.3	Cancerous prostate	76-34-1-C2-PU
ID236	est-not-ext	6.2	Hypertrophic prostate	77-8-1-F11-PU
ID237	est-not-ext	6.2	Cancerous prostate	76-41-1-F3-PU
ID238	est-not-ext	6.1	Cancerous prostate	76-22-3-G4-PU
ID239	est-not-ext	6.1	Normal prostate	78-40-1-A6-PU
ID240	est-not-ext	6	Normal prostate	78-41-2-H11-PU
ID241	est-not-ext	6	Normal prostate	78-6-3-A12-PU
ID242	est-not-ext	6	Hypertrophic prostate	77-25-1-A6-PU
ID243	est-not-ext	5.9	Hypertrophic prostate	77-35-2-E4-PU
ID244	est-not-ext	5.9	Hypertrophic prostate	77-36-1-G4-PU
ID245	est-not-ext	5.8	Hypertrophic prostate	77-40-3-D6-PU
ID246	est-not-ext	5.8	Normal prostate	78-17-3-A3-PU
ID247	est-not-ext	5.7	Normal prostate	78-33-3-D7-PU
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SEQ. ID	CATTCODY	VON HELINE	TISSUE	INTERNAL
<u>NO.</u>	<u>CATEGORY</u>	_SCORE	SOURCE	DESIGNATION
ID248	est-not-ext	5.7	**	
ID249	est-not-ext	5.7 5.7	Hypertrophic prostate	77-23-4-E10-PU
ID250	est-not-ext	5.7 5.7	Cancerous prostate	76-25-4-F11-PU
ID251	est-not-ext	5.7 5.7	Cancerous prostate	76-33-2-F8-PU
ID252	est-not-ext	5.7 5.7	Normal prostate	78-47-4-D6-PU
ID253	est-not-ext	5.6	Normal prostate Cancerous prostate	78-34-4-G6-PU
ID254	est-not-ext	5.6	Normal prostate	76-23-3-G8-PU
ID255	est-not-ext	5.6	Cancerous prostate	78-41-1-A6-PU
ID256	est-not-ext	5.5	Normal prostate	76-38-1-E4-PU
ID257	est-not-ext	5.4	Cancerous prostate	78-2-4-F11-PU
ID258	est-not-ext	5.4	Normal prostate	76-13-3-A9-PU
ID259	est-not-ext	5.2	Cancerous prostate	78-7-3-D9-PU
ID260	est-not-ext	5.1	Hypertrophic prostate	76-6-2-G5-PU
ID261	est-not-ext	5	Hypertrophic prostate	77-39-4-H4-PU
ID262	est-not-ext	5	Normal prostate	77-13-3-F1-PU
ID263	est-not-ext	4,9	Hypertrophic prostate	78-24-4-A4-PU
ID264	est-not-ext	4.9	Cancerous prostate	77-1-2-B4-PU
ID265	est-not-ext	4.9	Cancerous prostate	76-42-2-F3-PU 76-40-3-G6-PU
ID266	est-not-ext	4.8	Cancerous prostate	76-44-1-E3-PU
ID267	est-not-ext	4.8	Hypertrophic prostate	77-3-4-HI-PU
ID268	est-not-ext	4.8	Cancerous prostate	76-45-2-C4-PU
ID269	est-not-ext	4.8	Prostate	60-12-1-D7-PU
ID270	est-not-ext	4.8	Normal prostate	78-46-2-B4-PU
ID271	est-not-ext	4.7	Prostate	60-12-3-A7-PU
ID272	est-not-ext	4.7	Normal prostate	78-24-3-A8-PU
ID273	est-not-ext	4.6	Hypertrophic prostate	77-17-3-A7-PU
ID274	est-not-ext	4.6	Hypertrophic prostate	77-10-1-F6-PU
ID275 ID276	est-not-ext	4.5	Prostate	60-13-1-E11-PU
ID276 ID277	est-not-ext	4.4	Normal prostate	78-24-3-C6-PU
ID277	est-not-ext	4.4	Cancerous prostate	76-23-1-B4-PU
ID278	est-not-ext	4.3	Hypertrophic prostate	77-9-1-E2-PU
ID280	est-not-ext	4.2	Normal prostate	78-4-4-B10-PU
ID281	est-not-ext est-not-ext	4.2	Normal prostate	78-30-2-C1-PU
ID282	est-not-ext	4.2	Normal prostate	78-38-2-E9-PU
ID283	est-not-ext	4.2 4.1	Normal prostate	78-8-2-F2-PU
ID284	est-not-ext	4.1 4.1	Cancerous prostate	76-20-3-H1 -P U
ID285	est-not-ext	4.1 4.1	Cancerous prostate	76-14-1-B3-PU
ID286	est-not-ext		Normal prostate	78-18-4-D6-PU
ID287	est-not-ext	4 4	Hypertrophic prostate	77-11-4-B3-PU
ID288	est-not-ext	4	Normal prostate	78-16-2-C2-PU
ID289	est-not-ext	3.9	Hypertrophic prostate	77-38-2-G5-PU
ID290	est-not-ext	3.9	Normal prostate Hypertrophic prostate	78-25-1-H11-PU
ID291	est-not-ext	3.8	Cancerous prostate	77-12-3-H7-PU
ID292	est-not-ext	3.8	Normal prostate	76-21-4-A3-PU
ID293	est-not-ext	3.7	Cancerous prostate	78-41-1-C6-PU
ID294	est-not-ext	3.7	Cancerous prostate	76-5-2-H11-PU
ID295	est-not-ext	3.7	Cancerous prostate	76-8-4-D9-PU
ID296	est-not-ext	3.7	Prostate	76-18-2-D4-PU
ID297	est-not-ext	3.7	Hypertrophic prostate	60-12-3-G4-PU
ID298	est-not-ext	3.6	Cancerous prostate	77-20-2-E11-PU
ID299	est-not-ext	3.6	Normal prostate	76-1-2-G6-PU
ID300	est-not-ext	3.6	Normal prostate	78-8-3-F2-PU
			F- 400010	78-12-4-E9-PU

SEQ. ID NO.	CATEGORY	VON HEIJNE SCORE	TISSUE SOURCE	INTERNAL DESIGNATION
ID301	est-not-ext	3.6	Hypertrophic prostate	77-15-2-E2-PU
ID302	est-not-ext	3.5	Cancerous prostate	76-7-3-A12-PU
ID303	est-not-ext	3.5	Normal prostate	78-22-3-E10-PU
ID304	est-not-ext	3.5	Hypertrophic prostate	77-2-3-E11-PU
ID305	est-not-ext .	3.5	Normal prostate	78-29-1-B2-PU
ID306	ext-vrt-not-genomic	12	Normal prostate	78-47-2-C1-PU
ID307	ext-vrt-not-genomic	12	Normal prostate	78-43-4-G12-PU
ID308	ext-vrt-not-genomic	12	Hypertrophic prostate	77-38-1-A8-PU
ID309	ext-vrt-not-genomic	8.9	Normal prostate	78-45-4-F12-PU
ID310	ext-vrt-not-genomic	8.1	Normal prostate	78-35-3-D1-PU
ID311	ext-vrt-not-genomic	7.7	Normal prostate	78-10-1-H8-PU
ID312	ext-vrt-not-genomic	6.9	Cancerous prostate	76-43-1-E3-PU
ID313	ext-vrt-not-genomic	5.9	Normal prostate	78-29-2-C10-PU
ID314	ext-vrt-not-genomic	5.3	Hypertrophic prostate	77-38-3-B11-PU
ID315	ext-vrt-not-genomic	5.1	Normal prostate	78-36-4-A8-PU

TABLE III

	TABLE III
SEQ. ID	
NO.	SIGNAL PEPTIDE
<u></u>	4-Strad Lot Tibe
ID38	MVFVHLYLGNVLALLLFVHYSNG
ID39	MGMCFAAESDVQMFIAFLLCIFLICAALA
ID40	MAVRELCFSRQRQVLFLFLFWGVSLA
ID41	MRILQLILLALATGLVGG
ID42	MRILQLILLALATGLVGG
ID43	MRSCLWRCRHLSQGVQWSLLLAVLVFFLFA
ID44	MRILQXILLALATGLVGG
ID45	MLEECGAGVDLGFGGVKFASETPNLLWLLLKLVSTXWA
ID46	MIACSIRELHRCLLLALVAESSS
ID47	MGPPSLVLCLLSATVFS
ID48	MPGPRVWGKYLWRSPHSKGCPGAMWWLLLWGVLQX
ID49	MHRPEAMLLLITLALLGGPTWX
ID50	MVSVSLALLSGWVGS
ID51	MHIFSICCMXSELHKMKSLSLQLASEKRSLVALVEEIVFLLLRVSPCLG
ID52	MKLWVSALLMAWFGVLS
ID53	MKVLISSLLLLLPLMLMSMVSS
ID54	MKVLISSLLLLLPLMLMSMVSS
ID55	MLLLLQLSLPSPTS
ID56	MLKMLSFKLLLLAVALG
ID57	MHRPEAMLLLLTLALLGXXXWA
ID58	MLKVSAVLCVCAAAWC
ID59	MKVGVLWLISFFTFTDG
ID60	MCIILLDLICLLFITA
ID61	MDCASISVKFTSMATMHDLSQFWASRGEVTNWWPVGQTSLPLFYLAFMVFGSFFPLISC
ID62	MIASIDILAVERGIAGAIG
ID63	MVCVLVLAAAAGAVA
ID64	MKKTGDGGTLSTERIGGAALLSLLLKRMKMTLMIPLLLLTPITA
ID65	MELGCWTQLGLTFLQLLLISSLP
ID66	MRXKWKMGGMKYIFSLLFFLLLEGGXT
ID67	MRGATRVSIMLLLVTVSDC
ID68	MIAISAVSSALLFSLLCEAST
ID69	MIAISAVSSALLFSLLCEAST
ID70 -	MDPNGGCCTLLTLVLCVAVAYE
ID71	MEGEIYFQVFLSLFTFSTSLPSSLS
ID72	MYVVAMFGNCIVVFIVRTERSLHAPMYLFLCMLAAIDLALS
ID73	MRETXPLPKPLKDTAPSSHGVGSDSPSATRPWFLAPWCPGTOS
ID74	MDRPGSLSVFGSLPASLGTWLSSPAWLVDRPVRSAHPSANSTGVRMSVJ VAJ ALBSLCDS
ID75	MH I L A A CK A ILLER A PS CCCCC
ID76	MDLNSASTVVLQVLTQATS
ID77	MSSCNFTHATFVLIGIPGLEKAHFWVGFPLLSMYVVAMFGNCIVVFIVRTERSLHAPMYL
	FLCMLAAIDLALS
ID78	MYRLSLIAGPGSYPVLRWGVWDIPSSLVQVTYHQPNLTTNLDLPLFFSCSISATHS
ID79	WE VDGPSERPALCELLA VAMSFF
ID80	MPCSLTWRLPPRTCQXXGLXKSXLXXLLTPPPSYG
ID81	MVXWLVLFALQIYSYXSTRDQPASRXRLLFLFLTSIAEXCS
ID82	MARHGLPLLXXXSLPVGA
ID83	MVHLRTGLMLMSADRLRTLYYTVTILYILWYCSVCSS
ID84	MGILSTVTALTFARA
ID85	MELGCWTQLGLTFLQXLLISSLX
ID86	MELLRVCSFFLLCXSVFTDCKG
ID87	MIVRPRLNLTWFLLLPPGQCRA

SEQ. ID	
NO.	SIGNAL PEPTIDE
ID88	MQFLFKMVALCCCLWKISG
ID89	MLKVSAVLCVCAAAXXSQSLX
ID90	MSMQFLFKMVALCCCLWKISG
ID91	MAQHLWILLGSLSCRTS
ID92	MNKEXVSXERXAQVRLYLFSGFWTFXLG
ID93	MVLWRAKIXRNVPVTLSEENRSEGKVGFQAYKNYFRAGAHWIVFIFLILLNTAA
ID94	MLLXFFTSVLWLTSPSQP
ID95	MELISPTVIIILGCLALFLLLO
ID96	MHGFEIISLKEESPLGKVSQGPLFNVTSGSSSPVTWLGLLSFONLHC
ID97	MTWVRHAPGKSLEWVATVTDGGDKTFYAASVKGRFNVSRDNSKNTLFLHLSGLSAA
ID98	MLTSFFSLTANCQS
ID99	MLLCLLTPLFFMXPTGFS
ID100	MDDDYEAYHSLFLSLLGLCPS
ID101	MEWGKQWLVWLLLGHMVVS
ID102	MRRGKRLLESQSSSPKACLQLGFETELTQGVLWILVIQA
ID103	MVAATEAALLESVVWLPCHG
ID104	MSWNPSVSLPLLSSWGSTA
ID105	MKRIQGILFLILLSLHLERRWT
ID106	MVQRLWVSRLLRHRKAQLXLXNLLTFGLEVCLAAG
ID107	MAAGVPFALVTSCSSVFS
ID108	MTVFLXFCFPRCHS
ID109	MXPNNFWQKLGRKKPRIFTCTQSSTGEAAVKAENLILLEVFVWNGLQG
ID110	MFRSDRMWXCHWKWKPSPLLFLFALYIMCVPHSVWG
ID111	MTQRSIAGPICNLKFVTLLVALSSELPFLGA
ID112	MIPLLLLRSACN
ID113	MXSPLPVLLLSXNLNLIIQ
ID114	MLMCKMLKSQKNCQENXXIKIILFLKPMCSPQYLLTFLVFTXKLSS
ID115	MKKKSSPNQYLHSSLHXIRLFSFLHFSEEGVLLLAIDLKIIVILHCAASIIS
ID116	MFSCFFSTSLATSVSLEAQSCFA
ID117	MHHGLTPLLLGVHEQKQQVVKFLIKKKANLNALDRYGRTALILAVCCGSA
ID118	MSPCIYFFACFQALTSS
ID119	MAEEMESSLEAXFSSSGAVSGASGFLPPARS
ID120	MAEEMESSLEASFSSSGAVSGASGFLPPARS
ID121	MLVLGSPLLGPLLWHLSLILLKPLCLP
ID122	MHLLDLESMGKSSDGKSYVITGSWNPKSPHFQVVNEETPKDKVLFMTTAVDLVIT
ID123	MENLKDFYVLFVFSSIPLTFL
ID124	MPQYCLSIFSLVLPVCRM
ID125	MVAPVLETSHVFCCPNRVRGVLNWSSGPRGLLAFGTSCSVVLY
ID126	MPIIDQVNPELHDFMQSAEVGTIFALSWLITWFGHXLS
ID127	METXCPCCCCPCXGXGSLXXKPVYELQVQKSVTVQEGLCVLVPCSXSXX
ID128	MSPCTYFFACFXXLTSS
ID129	MGRGERRHYWGPKLVLKCLSFSXPSLP
ID130	MSQDGGXGELKHMVMSFRVSELQVLLGFAGRNKSGRKHELLAKALHLLKSSC
ID131	MHHRMNEMNLSPVGMEQLTSSSVSNALPVSGSHLGLAASPTHSAIPAPGLPVAIPNLGPS
	LSSLPSALS
ID132	MLHSDNIWNLFSLFSTSTT
ID133	MQPASPPARWSFHSAAGWSGGQA
ID134	MCFSFLLAGSISHMFSQA
ID135	MYGFIIGLSILFHCSVCLFLC
ID136	MSFGXILTFRVSLLGCXLAININT
ID137	MAVYVGMLRLGRLCAGSSGVXG
ID138	MFNTIYLVISLVSIFFFWEVTNA
ID 139	MALPPKGCGSLPLTTGSSWSLS

SEQ. ID	•
NO.	SIGNAL PEPITDE
_1 4 O	SIGNAL PEPTIDE
ID140	MFVFLSWASFLAPLLR
D 141	MXMKSANKITLLXHHLLSCSPLXPLGKS
ID142	MCNYNIYVLYNIGYLYHPKSFLLLFIVIPQTP
ID143	MAVAMVKLCERAGLPLLAAPLLRSLLP
ID144	MINION DEVOLUE AT SUCCESSION OF THE SECOND O
22144	MLNVVRALRXPQWCAEYCLSIHYQHGGVICTQVHKQTVVQLALRVADEMDVNIGHEVGYV
ID145	TIPLE COLLECTION OF THE CONTRACT OF THE COLLECTION OF THE COLLECTI
ID146	MINOUENASAQNALAGECIGS I SY VHG
ID147	MLNGPFQHRNSRIMTHRSAEKTLLGSLSLWRWSAM
ID148	MRVKDPTKALPEKAKRSKRPTVPHDEDSSDDIAVGLTCQHVSHA
ID148 ID149	MPQKGLGLLGILSGDFSLLALSMLKGTG
ID149	MAMWNRPXXXLPQQPLXAEPTAEGEPHLPTGRXXTEANRFAYAALCGISLSQLFP
ID150	MECLODELESE MAIABAMS
ID151	MQENAHNLRLFKCLLIYFLGLAADTYF
ID152 ID153	MHTCSLPCLLFAQLLEFCSFPPDVPHNCAPIVSVRPPNIVAAFEGCSVATALFPPLCIS
ID153	MQQROAAGSRGCALFFLL(IVLFF()(IVV)
ID154	MXXSIFISEKYGLCPSKTPIMKMLPSLILNRSLPTASSS
ID156	MAFDVSCFFWVVLFSAGCKV
1130	MEVAANCSLR VKRPLLDPRFEGYKXSLEPLPCYQLELDAAVAXVKLRDDQYTLEHMHAFG
ID157	MINITEREDS WIQDSVIIIDILGRIMNLIVMLITAXG
ID158	MNVGTAHXXVNPNTRVMNSRGIWLSYVLAIGLLHIVLLS MENFNMYKNKSWWTLLSSSPSFM
ID150	MANACTARICE A DISTRICT OF STATE OF STAT
ID160	MNVGTXHSEVNPNTRVMNSRGIWLSYVLAIGLLHIVLLS MAAASAVSVLLVAA
D161	
ID162	MAYSKASGSPVLSQAVPGENASHRRGSADLGSGSGLSWARLSQS
ID163	MKPRRNLEEDDYLHKDTGETSMLKRPVLLHLHQTAHA MIICYDIPCAHMLVCPTIG
ID164	MYSSEDSTLASVPPAATFG
ID165	MCEDDAODDANAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
103	MGEDPXQPRKYKKXKXELQGDXPPSSPTNDPTVKYETQPRFITATGGTLHMYQLEGLNWL RFSWA
ID166	IGUIA
ID167	MFYVAMTKTHKRIRSLCNIHHGLFQFTQQLLGCLQCCWLQSG
ID168	MVSPKDLPLVLLQDIKVPSSMTGSHAGNPHIERNDLPRHGSPQFFTGXTCASXNPSQCLA MEFXSLFCLYFSCFL
ID169	
ID170	MALHFQSLAELEXLCTHLYIGTDLTQRIEAEKALLELIDSPECLS MRTLFGAVRAPFSSLTLLLITPSPSPL
ID171	MBHSL I KCISAONS AADVVDAOL MATALISON LANGUAGE
ID172	MRHSLLKGISAQIVSAADKVDAGLPTAIAVSSLIAVGTSHG MTLSCFIFFYISSLC
ID173	MILCFLLPHHRLQEA
ID174	MFSLFALNMPLGFC
D175	MASSPGVAMHSLWATIHTSVWGVLPPPACSA
D176	MSQEGAVPASAVPLEELSSWPEELCRRELPSVLPRLLSLSQHSES
D177	MTRECPSPAPGPGAPLSGSVLAEAAVVFAVVLSIHA
D178	MQELHLLWWALLIGLAQA
D179	MGRQALLLLALCATGAQG
D180	MGPSTPLLILFLLSWSGPLQG
D181	MSCRELTHRPCSPHLLLLCPLSRGCCP
D182	MGWTMRLVTAALLLGLMMVVTG
D183	MKFLIFAFFGGVHLLSLCSGKVYA
D184	MQCFSFIKTMMILFNLLIFLCGAALLAVG
D185	MWAFSELPMPLLINLIVSLLGFVATVTL
D186	MASSNTVLMRLVASAYSIA
D187	MKFLIFAFFGGVHLLSLCSGKAIC
D188	MADTTPNGPQGAGAVQFMMTNKLDTAMWLSRLFTVYCSALXVLPLLGLHEA
D189	MRFRHFXKXIGXVLVLSVVXXAMA

CEO ID	
SEQ. ID	SIGNAL DEPEND
<u>NO.</u>	SIGNAL PEPTIDE
ID190	MELGSCLEGGREAAEEEGEPEVKKRRLLCVEFASVASCDA
D191	MASPFSGALQLTDLDDFIGPSQECIKPVKVEKRAGSGVAKIRIEDDGSYFQINQDGXTRRLE
13171	KAKVSLNYCXACSGCITSAETVLITQQSHEELKKVLDANKMAAPSQQRLVVVSVSPQSRA
ID192	MGPVPTAVAGAGSRLVKPSQTLSLTCAVSGGSLVAELLLGAGSG
ID193	MESGGRPSLCQFILLGTTSVVTA
ID194	MQVCRCIYIICFXLPPLFS
ID195	MAQRLLLRFLASVIS
ID196	MLFIFNFLFSPLPTPALICILTFGAAIFLWLITRPQPVLP
ID197	MYPKWEAPVTFCQLKREKDPPHPAHSPFLQPRFSHMLQLLPSKALC
ID198	MALYQRWRCLRLQGLQACRLHTAVVSTPPRWLAERLGLFEELWA
ID199	MGVPRPQPWAXGLLLFLLPGSLG
ID200	MAAAVPKRMRGPAQAKLLPGSAIQALVGLARPLVLALXLVSAALS
ID201	MWLWEDQGGLLGPFSFLLLVLLLVTRXRS
ID202	MNWELLLWLLVLCALLLLLVQLLRFLRA
ID203	MEKIPVSAFLLLVALSYTLA
ID204	MSNYTDAESSFSKQEIIRVAMEKIPVSAFLLLVALSYTLA
ID205	MQFXTWATSSSQPALWSLLLVSWAAMVLRLRSKCALVTFFFILLLIFIAEVAA
ID206	MNWELLLWLLVLCALLLLLVHLLRFLRA
ID207	MTTFLPVPQMMAGFSFGTFGNPPMESPSAWQTIHQPFIVSCLTLWSPGCWP
ID208	MASKGMRHFCLISEQLVXFSLLATAILG
ID209	MAAAAWLQVLPVILLLLGAHP
ID210	MASPRTVTIVALSVALGLFFVFMGTIKLTPRLSKDAYSEMKRAXKSYVRALPLLKKMGIN
ID211	SILLRKSIGALEVACGIVMTLVPGRPKDVANFFLLLLVLAVLFFHQLVG
ID211 ID212	MPNLSFGGLDTNQMRVNFLSVDVCKLLLLCALHSHTYC
ID212 ID213	MGPPMLQEISNLFLILLMMGAIFTLAALKESLSTCIPAIVCLXXLLLLNVGQLLA
110213	MXXFTDPSSVNEKKRREREERQNIVLWRQPLITLQYFSLEILVILKEWTSKLWHRXXIVV XFLLLAXLIA
ID214	MPLLRGLLWXQVLCA
ID215	MKLLSLVAVVGCLLVPPAEA
ID216	MPALLPVASRLLLLPRVLLTMASG
ID217	MCLLLGATGVGKTLLVKRLQEVSSRDGKGDLGEPPPTRPTVGTNLTDIVAQRKITIRELG
	GCMGPIWSSYYGNCRSLLFVMDASDPTQLSAXXVQLLGLLSAEQLAEA
ID218	MELPAVNLESDSPRSLAADNLGLHCILRLLCLGQLHHPGLG
ID219	MAFLRKVYSILSLQVLLTTVTSTVFLYFESVRTFVHESPALILLFALGSLG
ID220 -	MYTYGNKQHNSPTWDDPTLAIALAANAWA
ID221	MQQIFIQQCRELNFWSREPWILVLALPLTVWP
ID222	MKAVLLALLMAGLAL
ID223	MGLQACLLGLFALILS
ID224	MRPGQVSLLGPDAVSVLGSGLGLSPGTSS
ID225	MINPSVPSKSNSHPFLSTVMFTSASLLLPMSTG
ID226	MSEKEXNFPPLPKFIPVKPCFYQNFSDEIPVEHQVLVKRIYRLWMFYCATLGVNLIACLA
W000	WWIGGGSG
ID227	MNPTKLILKTILRLYFFLQLAHS
ID228	MASSSPDSPCSXXCFVSVPPASA
ID229	MXPVLAALAHVLCPYMAPGLCREPIRXLIAXLEPPGAMA
ID230	MNNLNDPPNWNIRPNSRADGGDGSRWNYALLVPMLGLAAFRWIWS
ID231 ID232	MLLFLAALCSLFFFLSLQ MLFLGKVLIVCSTGLAGIMLLNVOODVTNUURRING DLUVGLEAFLAVAGE
ID232 ID233	MLFLGKVLIVCSTGLAGIMLLNYQQDYTVWVLPLIIVCLFAFLVAHC
ID233	MQGIPILTPVTTQSIAISIVLTVQGLLLLVHSFWFTVC MQNFCHHLAICTVILFCVLLSLRPHTS
ID234 ID235	MPSFSKDLLTVPKLGTGHXXGXGSYDXALXLLLKCLWSNVVPECTMASSNTVLMRLVASA
	YSIA YSIA
ID236	MRGAHLTALEMLXAFASHIXA
	· · · · · · · · · · · · · · · · · · ·

SEQ. ID	
NO.	SIGNAL PEPTIDE
ID237	MEVGI DA ITI EL TCA CODUCIATION CONTRACTOR DE CONTRACTOR D
ID 257	MEVGLPAITLFLTSASSPVVATTMDQEPVGGVERGEAVAASGXAAAAAFGESAGQMSNER
	GFENVELGVIGKKKKVPRRVIHFVSGETMEEYSTDEDXVDGLEKXMFCLLLIRQNLPGVP TYGFTCFGLLHQLSQCVTS
ID238	MKELERQQKEVEERPEKDFTEKGSRNMPGLSAATLASLGGTSS
ID239	MSMGFMMLVLVILCIVTVCVT
ID240	MMELXLKXXTKXEXESACTEAYSQSDEQYACHLGCQNQLPFAELRQEQLMSLMPKMHLLF
	12.27101 110
ID241	MVSNASETSCLGLILLFASHLINQ
ID242	MPRKRKCDLRAVRVGLLLGGGGVYGSRERETEPGCPALSDWDVDVDVDDDGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAG
	EIVI ID VCI ALLA
ID243	MGMWSIGAGALGAAALALLLANT
ID244	MDVAFLEXLIKDDIERGRLPLLLVANAGTAA
ID245	MRTLFNLLWLALACSP
ID246	MNAQPGLXLDCITRFLTXGQFICLQWALPHSEA
ID247	MGKEWGWQEMENGGAAPAWGAGPPVHPAPPPVEKTI.SWGCGFGI.HSGFGGSGCGVGI.GD
TD 0.40	DCD (100) CC
ID248 ID249	MAAPSGGWNGVGASLWAALLLTATVRLSA
ID249 ID250	MIAIYGKNFCVSAKNAFMLLMRNIVRVVVLDKVIDLLLFFGKLLVVGG
110230	WERNCRUST GVIREGUIDIXETK ARRTVWFPR GDVCFD VTDDDA VDVTG GGC A DD A VDV
ID251	DAGATOL LACELLINE A LUALINA VICE IN
10231	MFVEYRKQLKLLLDRLAQVSPELLLASVRRVFSSTLQNWQTTRFMEVEVAIRLLYMLAEA
ID252	MLLGTSNIIIFLIQWHGSVFQ
ID253	MXNRFATAFVXACVLSLIST
ID254	MSLTSGFLRVSQG
ID255	MANFKGHALPGSFFLIIGLCWSVKYPLKYFSHTRKNSPLHYYQRLEIVEAAIRTLFSVTGILA
ID256	MQDTGSVVPLHWFGFGYAALVASGGIIGYVKAGSVPSLAAGLLFGSLAGLGA
ID257	MEXGLKSADPRDGTGYTXXXXYCCALLTSLXCIWG
ID258	MASPSRRLOTKPVITCFKSVLLIVTFIFWTGVILLAVGIVG
ID259	MESKELAPTRIGGASSGSRSGGTLISTAPI.TTRVI NPTAOCECI DCTI PRIMOTIR SVELI
	I CHOKWD
ID260	MSMAVETFGFFMATVGLLMLGVTLPNSYW
ID261	MEKIPVSXFLXLXXLSXXWP
ID262	MHSAEEPLXLAALRGARGHLPCGSRHHVGSLAPASVPAPGACLWVCEWETLLPGLILERP
ID263	LYFSALA
ID263 ID264	MAGQFRSYVWDPLLILSQIVLMQTVYYGSLGLWLALVDGLVRX
11)204	MAPKVFRQYWDIPDGTDCHRKAYSTTSIASVAGLTAAAYRVTLNPPGTFLEGVAKVGQYT FTAAAVGAVFGLTTCISA
ID265	MAAAAWLQVLPVILLLLG
ID266	MEIYFIFCIIVPIAAATVYKSWCLLLILDMNVLYTDA
ID267	MSRYTSPVNPAVFPHLTVVLLAIGMFFTAWF
ID268	MRLAAEAHPGRTHTLFRRLKPFLMLSSSLPLLIWL
ID269	MLEHLXSLPTQMDYKGQKLAXQMFQGIILFSAIVGFIYG
ID270	MEYSKVLFCSFSNVLG
ID271	MASKIGSRRWMLQLIMQLGSVLLTRC
ID272	MEHYRKAGSVELPAPSPMPOLPPDTLEMRVRDGSKIRNI I GLALGPLEGGSA
ID273	WINALWIVLENVIVVLIALTCLDGTTVS
ID274	MNWSIFEGLLSGVNKYSTAFGRIWLSLVFIFRVLVYLVTAERVWS
ID275	MIISLEIYIEXTCSNT
ID276	MFRLNSLSALAELAVG
ID277	MTAGTLRTWLCCAGSWA
ID278 ID279	MLGRPCFHSPQRLLVILCVSVKAG
11/4/7	MDEARDNACNDMGKMLQFVLPVATQIQQ

SEQ. ID	
NO.	SIGNAL PEPTIDE
ID280	MSPISIRELCALGSAPSSMWA
ID281	MTDLLSASPWALT
ID282	MSWSGLLHGLNTSLTCGPALVPRLWA
ID283	MADVINVSVNLEAFSQAISAIQA
ID284	MNVIDHVRDMAAAGLHSNVRLLSSLLLTMSNN
ID285	MTSACLAWTAVRPSAC
ID286	MNGSRTLTHSISDGQLQGGQSNSELFQQEXQTAPAQVPQGFNVFGMSSSSGASNS
ID287	MLGFFLFLSFVLMYDG
ID288	MMEERANLMHMMKLSIKVLLQSALSLG
ID289	MELEXIVSAALLAFVOT
ID290	MLRQIIGQAKKHPSLIPLFXFIGTGA
ID291	MVKETQYYDILGVKPSASPERSRRPIGSWRSSTTRTRTRMRARSLNSYPRHMKCFQIQRK
	GMFMTKAESRQXKKEAQAAPASLHPWTSLTCSLVVVDG
ID292	MANLFIRKMVNPLLYLSRHTVKPRALSTXLFGSIRG
ID293	MAAAAASRGXGAKLGLRXIRIHLCQRSPGSQG
ID294	MFPSCYLCYSLCGSILLSIFSAYNRLSLMLRIALTLIPSMLSRA
ID295	MSTQXGLSMHAHPQAYTPFIYLHARKRRGEIGDADSRFNDRYAHKSAQLXFLYFVCCIFQ
ID296	MKHFQDLPSSCSCSLISFTRG
ID297	MSQRSLCMDTSLDVYRXLIELNYLGTVSLTKCVLPHMIERKXXKIVTVNSILGIISVPLSIG
ID298	MGGSGSRLSKELLAEYQDLTFLTKQEILLAHRRFCELLPQEQRXXSRHFGHKCPSSRFSA
	FQSSRPTPSRSESAGSSPHPQPKTALALRTSWISSVCS
ID299	MWRLLARASAPLLRVPLSDSWALLPASA
ID300	MADHVQSLAQLENLCKQLYETTDTXXRSSXAEKALVEFTNSPDCLSKCQLLLERGSSSYS
	QLLAATCLTKLVSRTNNPLPLEQRIDIRNYVLNXLATRPKLATFVTQALIQXYA
ID301	MAYHGLTVPLIVMSVFWGFVGFLVPWFIPKGPNRGVIITMLVTCSVCCYLFWLIA
ID302	MSTGQLYRMEDIGRFHSQQPGSLTPSSPTVGEIIYNNTRNTLGWIGGILMGSFQGTIA
ID303	MGWQRWWCFHLQAEASA
ID304	MSVIFFACVVRVRDG
ID305	MAVTALAAXTWLGVWG
ID306	MSLSAFTLFLALIGGTSG
ID307	MSLSAFTLFLALIGGTSG
ID308	MSLSAFTLFLALIGGTSG
ID309	MVELMFPLLLLLPFLLYMA
ID310	MWLLYLLVPALFCRA
ID311	MKQILHPALETTAMTLFPVLLFLVAGLLPSFP
ID312	MLKALFLTMLTLALVKS
ID313	MEKNPLAAPLLILWFHLDCVSS
ID314	MRVVTIVILLCFCKA
ID315	MDQFPESVTENFEYDDLAEACYIGDIVVFGTVFLSIFYSVIFAIGLVGNLLVVFALTNSK
	KPKSVTDIYLLNLALSDLLFVATLPFWTHY

Minimum signal peptide score	false positive rate	false negative rate	proba(0.1)	proba(0.2)
3.5	0.121	0.036	0.467	0.664
4	0.096	0.06	0.519	0.708
4.5	0.078	0.079	0.565	0.745
5	0.062	0.098	0.615	0.782
5.5	0.05	0.127	0.659	0.813
6	0.04	0.163	0.694	0.836
6.5	0.033	0.202	0.725	0.855
7	0.025	0.248	0.763	0.878
7.5	0.021	0.304	0.78	0.889
8	0.015	0.368	0.816	0.909
8.5	0.012	0.418	0.836	0.92
9	0.009	0.512	0.856	0.93
9.5	0.007	0.581	0.863	0.934
10	0.006	0.679	0.835	0.919

TABLE IV

Minimum signal peptide score	All ESTs	New ESTs	ESTs matching public EST closer than 40 bp from beginning	ESTs extending known mRNA more than 40 bp	ESTs extending public EST more than 40 bp
3.5	2674	947	599	23	150
4	2278	784	499	23	126
4.5	1943	647	425	22	112
5	1657	523	353	21	96
5.5	1417	419	307	19	80
6	1190	340	238	18	68
6.5	1035	280	186	18	60
7	893	219	161	15	48
7.5	753	173	132	12	36
8	636	133	101	11	29
8.5	543	104	83	8	26
9	456	81	63	6	24
9.5	364	57	48	6	18
10	303	47	35	6	15

TABLE V

Tissue	All ESTs	New ESTs	ESTs matching public EST closer than 40 bp from beginning	ESTs extending known mRNA more than 40 bp	ESTs extending public EST more than 40 bp
Brain	329	131	75	3	24
Cancerous prostate	134	40	37	1	6
Cerebellum	17	9	1	0	6
Colon	21	11	4	0	0
Dystrophic muscle	41	18	8	0	- 1
Fetal brain	70	37	16	0	- 1
Fetal kidney	227	116	46	1	19
Fetal liver	13	7	2	0	o
Heart	30	15	7	0	1
Hypertrophic prostate	86	23	22	2	2
Kidney	10	7	3	0	2
Large intestine	21	8	4	0	1
Liver	23	9	6	0	o
Lung	24	12	4	0	1
Lung (cells)	57	38	6	0	4
Lymph ganglia	163	60	23	2	12
Lymphocytes	23	6	4	0	2
Muscle	33	16	6	0	4
Normal prostate	181	61	45	7	11
Ovary	90	57	12	1	2
Pancreas	48	11	6	0	1
Placenta	24	5	1	0	o
Prostate	34	16	4	. 0	. 2
Spleen	56	28	10	0	1
Substantia nigra	108	47	27	1	6
Surrenals	15	3	3	1	0
Testis	131	68	25	1	8
Thyroid	17	8	2	0	2
Umbilical cord	55	17	12	1	3
Uterus	28	15	3	0	2
Non tissue-specific	568	48	177	2	
Total	2677	947	601	23	150

TABLE VI

129/4

Description of Transcription Factor Binding Sites present on promoters isolated from SignalTag sequences Promoter sequence P13H2 (546 bp):

Matrix	Position	Orientation	Score	Length	Sequence
CMYB_01	-502	+	0.983	. 9	TGTCAGTTG
MYOD_Q6	-501	•	0.961	10	CCCAACTGAC
S8_01	-444	•	0.960	11	AATAGAATTAG
S8_01	-425	+	0.966	11	AACTAAATTAG
DELTAEF1_01	-390	•	0.960	11	GCACACCTCAG
GATA_C	-364	•	0.964	11	AGATAAATCCA
CMYB_01	-349	+	0.958	9	CTTCAGTTG
GATA1_02	-343	+	0.959	14	TTGTAGATAGGACA
GATA_C	-339	•	0.953	11	AGATAGGACAT
TAL1ALPHAE47_01	-235	•	0.973	16	CATAACAGATGGTAAG
TAL1BETAE47_01	-235	+	0.983	16	CATAACAGATGGTAAG
TAL1BETAITF2_01	-235	+	0.978	16	CATAACAGATGGTAAG
MYOD_Q6	-232	•	0.954	10	ACCATCTGTT
GATA1_04	-217	• .	0.953	13	TCAAGATAAAGTA
IK1_01	-126	+	0.963	13	AGTTGGGAATTCC
IK2_01	-126	+	0.985	12	AGTTGGGAATTC
CREL_01	-123	+	0.962	10	TGGGAATTCC
GATA1_02	-96	+	0.950	14	TCAGTGATATGGCA
SRY_02	-41	•	0.951	12	TAAAACAAAACA
E2F_02	-33	+	0.957	8	TTTAGCGC
MZF1_01	-5	•	0.975	8	TGAGGGGA

Promoter sequence P16B4 (861bp):

Matrix	Position	Orientation	Score	Length	Sequence
NFY_Q8	-748	•	0.956	11	GGACCAATCAT
MZF1_01	-738	+	0.962	8	CCTGGGGA
CMYB_01	-684	+	0.994	9	TGACCGTTG
VMYB_02	-682	•	0.985	9	TCCAACGGT
STAT_01	-673	+	0.968	9	TTCCTGGAA
STAT_01	-673	•	0.951	9	TTCCAGGAA
MZF1_01	-556	•	0.956	8	TTGGGGGA
IK2_01	-451	+	0.965	12	GAATGGGATTTC
MZF1_01	-424	+	0.986	8	AGAGGGGA
SRY_02	-398	•	0.955	12	GAAAACAAAACA
MZF1_01	-216	+	0.960	8	GAAGGGGA
MYOD_Q6	-190	+	0.981	10	AGCATCTGCC
DELTAEF1_01	,-176	+	0.958	11	TCCCACCTTCC
S8_01	5	-	0.992	11	GAGGCAATTAT
MZF1_01	16	-	0.986	8	AGAGGGGA

Promoter sequence P29B6 (555 bp):

Matrix	Position	Orientation	Score	Length	Sequence
ARNT_01	-311	+	0.964	16	GGACTCACGTGCTGCT
NMYC_01	-309	+	0.965	12	ACTCACGTGCTG
USF_01	-309	+	0.985	12	ACTCACGTGCTG
USF_01	-309	•	0.985	12	CAGCACGTGAGT
NMYC_01	-309	•	0.956	12	CAGCACGTGAGT
MYCMAX_02	-309	•	0.972	12	CAGCACGTGAGT
USF_C	-307	+	0.997	8	TCACGTGC
USF_C	-307	•	0.991	8	GCACGTGA
MZF1_01	-292	•	0.968	8	CATGGGGA
ELK1_02	-105	+	0,963	14	CTCTCCGGAAGCCT
CETS1P54_01	-102	+	0.974	10	TCCGGAAGCC
AP1_Q4	-42	•	0.963	11	AGTGACTGAAC
AP1FJ_Q2	-42	•	0.961	11	AGTGACTGAAC
PADS_C	45	+	1.000	9	TGTGGTCTC

TABLE VII

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CLAIMS

- A purified or isolated nucleic acid comprising the sequence of one of SEQ ID
 NOs: 38-315 or comprising a sequence complementary thereto.
 - 2. The nucleic acid of Claim 1, wherein said nucleic acid is recombinant.
- 3. A purified or isolated nucleic acid comprising at least 10 consecutive bases of the sequence of one of SEQ ID NOs: 38-315 or one of the sequences complementary thereto.
- 4. A purified or isolated nucleic acid comprising at least 15 consecutive bases of one of the sequences of SEQ ID NOs: 38-315 or one of the sequences complementary thereto.
 - 5. The nucleic acid of Claim 4, wherein said nucleic acid is recombinant.
 - 6. A purified or isolated nucleic acid of at least 15 bases capable of hybridizing under stringent conditions to the sequence of one of SEQ ID NOs: 38-315 or one of the sequences complementary to the sequences of SEQ ID NOs: 38-315.
 - 7. The nucleic acid of Claim 6, wherein said nucleic acid is recombinant.
 - 8. A purified or isolated nucleic acid encoding a human gene product, said human gene product having a sequence partially encoded by one of the sequences of SEQ ID NO: 38-315.
- 20 9. A purified or isolated nucleic acid having the sequence of one of SEQ ID NOs: 38-315 or having a sequence complementary thereto.
 - A purified or isolated nucleic acid comprising the nucleotides of one of SEQ
 NOs: 38-315 which encode a signal peptide.
 - 11. A purified or isolated polypeptides comprising a signal peptide encoded by one of the sequences of SEQ ID NOs: 38-315.
 - 12. A vector encoding a fusion protein comprising a polypeptide and a signal peptide, said vector comprising a first nucleic acid encoding a signal peptide encoded by one of the sequences of SEQ ID NOs: 38-315 operably linked to a second nucleic acid encoding a polypeptide.
- 30 13. A method of directing the extracellular secretion of a polypeptide or the insertion of a polypetide into the membrane comprising the steps of:

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obtaining a vector according to Claim 12; and

introducing said vector into a host cell such that said fusion protein is secreted into the extracellular environment of said host cell or inserted into the membrane of said host cell.

- 14. A method of importing a polypeptide into a cell comprising contacting said cell with a fusion protein comprising a signal peptide encoded by one of the sequences of SEQ ID NOs: 38-315 operably linked to said polypeptide.
- 15. A method of making a cDNA encoding a human secretory protein that is partially encoded by one of SEQ ID NOs 38-315, comprising the steps of:

obtaining a cDNA comprising one of the sequences of SEQ ID NOs: 38-315;

contacting said cDNA with a detectable probe comprising at least 15 consecutive nucleotides of said sequence of SEQ ID NO: 38-315 or a sequence complementary thereto under conditions which permit said probe to hybridize to said cDNA;

identifying a cDNA which hybridizes to said detectable probe; and isolating said cDNA which hybridizes to said probe.

- 15 An isolated or purified cDNA encoding a human secretory protein, said human secretory protein comprising the protein encoded by one of SEQ ID NOs 38-315 or a fragment thereof of at least 10 amino acids, said cDNA being obtainable by the method of Claim 15.
 - 17. The cDNA of Claim 16 wherein said cDNA comprises the full protein coding sequence partially included in one of the sequences of SEQ ID NOs: 38-315.
 - 18. A method of making a cDNA comprising one of the sequences of SEQ ID NOs: 38-315, comprising the steps of:

contacting a collection of mRNA molecules from human cells with a first primer capable of hybridizing to the polyA tail of said mRNA;

25 hybridizing said first primer to said polyA tail;

reverse transcribing said mRNA to make a first cDNA strand;

making a second cDNA strand complementary to said first cDNA strand using at least one primer comprising at least 15 nucleotides of one of the sequences of SEQ ID NOs 38-315; and

isolating the resulting cDNA comprising said first cDNA strand and said second cDNA strand.

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- 19. An isolated or purified cDNA encoding a human secretory protein, said human secretory protein comprising the protein encoded by one of SEQ ID NOs 38-315 or a fragment thereof of at least 10 amino acids, said cDNA being obtainable by the method of Claim 18.
- 5 20. The cDNA of Claim 19 wherein said cDNA comprises the full protein coding sequence partially included in one of the sequences of SEQ ID NOs: 38-315.
 - 21. The method of Claim 18, wherein the second cDNA strand is made by:
 contacting said first cDNA strand with a first pair of primers, said first pair of primers
 comprising a second primer comprising at least 15 consecutive nucleotides of one of the
 sequences of SEQ ID NOs 38-315 and a third primer having a sequence therein which is
 included within the sequence of said first primer;

performing a first polymerase chain reaction with said first pair of nested primers to generate a first PCR product;

contacting said first PCR product with a second pair of primers, said second pair of primers comprising a fourth primer, said fourth primer comprising at least 15 consecutive nucleotides of said sequence of one of SEQ ID NO:s 38-315, and a fifth primer, said fourth and fifth primers being capable of hybridizing to sequences within said first PCR product; and

performing a second polymerase chain reaction, thereby generating a second PCR product.

- 20 22. An isolated or purified cDNA encoding a human secretory protein, said human secretory protein comprising the protein encoded by one of SEQ ID NOs 38-315, or a fragment thereof of at least 10 amino acids, said cDNA being obtainable by the method of Claim 21.
- The cDNA of Claim 22 wherein said cDNA comprises the full protein coding
 sequence partially included in one of the sequences of SEQ ID NOs: 38-315.
 - 24. The method of Claim 18 wherein the second cDNA strand is made by: contacting said first cDNA strand with a second primer comprising at least 15 consecutive nucleotides of the sequences of SEQ ID NOs: 38-315;

hybridizing said second primer to said first strand cDNA; and extending said hybridized second primer to generate said second cDNA strand.

- 25. An isolated or purified cDNA encoding a human secretory protein, said human secretory protein comprising the protein partially encoded by one of SEQ ID NOs 38-315 or comprising a fragment thereof of at least 10 amino acids, said cDNA being obtainable by the method of Claim 24.
- 5 26. The cDNA of Claim 25, wherein said cDNA comprises the full protein coding sequence partially included in of one of the sequences of SEQ ID NOs: 38-315.
 - 27. A method of making a protein comprising one of the sequences of SEQ ID NO: 316-593, comprising the steps of:

obtaining a cDNA encoding the full protein sequence partially included in one of the sequences of sequence of SEQ ID NO: 38-315;

inserting said cDNA in an expression vector such that said cDNA is operably linked to a promoter;

introducing said expression vector into a host cell whereby said host cell produces the protein encoded by said cDNA; and

- 15 isolating said protein.
 - 28. An isolated protein obtainable by the method of Claim 27.
 - 29. A method of obtaining a promoter DNA comprising the steps of:

obtaining DNAs located upstream of the nucleic acids of SEQ ID NO: 38-315 or the sequences complementary thereto;

screening said upstream DNAs to identify a promoter capable of directing transcription initiation; and

isolating said DNA comprising said identified promoter.

- 30. The method of Claim 29, wherein said obtaining step comprises chromosome walking from said nucleic acids of SEQ ID NO: 38-315 or sequences complementary thereto.
- 25 31. The method of Claim 30, wherein said screening step comprises inserting said upstream sequences into a promoter reporter vector.
 - 32. The method of Claim 30, wherein said screening step comprises identifying motifs in said upstream DNAs which are transcription factor binding sites or transcription start sites.
- 30 33. An isolated promoter obtainable by the method of Claim 32.

- 34. An isolated or purified protein comprising one of the sequences of SEQ ID NO: 316-593.
- 35. In an array of discrete ESTs or fragments thereof of at least 15 nucleotides in length, the improvement comprising inclusion in said array of at least one of the sequences of SEQ ID NOs: 38-315, or one of the sequences complementary to the sequences of SEQ ID NOs: 38-315, or a fragment thereof of at least 15 consecutive nucleotides.
- 36. The array of Claim 35 including therein at least two of the sequences of SEQ ID NOs: 38-315, the sequences complementary to the sequences of SEQ ID NOs: 38-315, or fragments thereof of at least 15 consecutive nucleotides.
- The array of Claim 35 including therein at least five of the sequences of SEQ ID NOs: 38-315, the sequences complementary to the sequences of SEQ ID NOs: 38-315, or fragments thereof of at least 15 consecutive nucleotides.

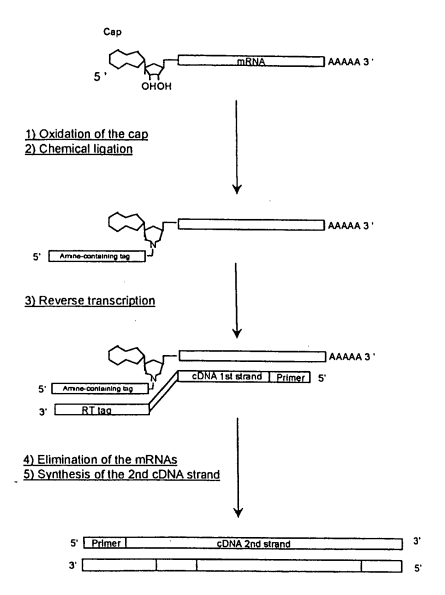


Figure 1

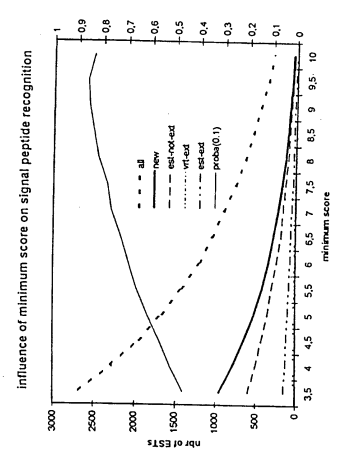


Figure 2

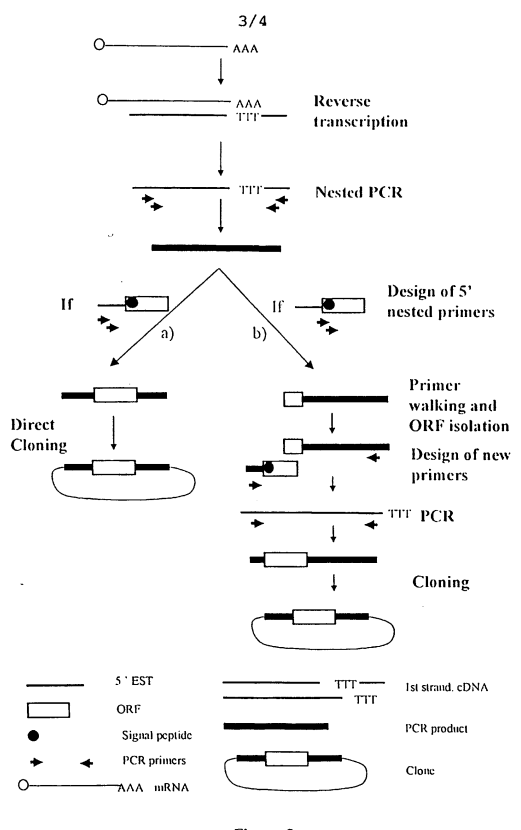
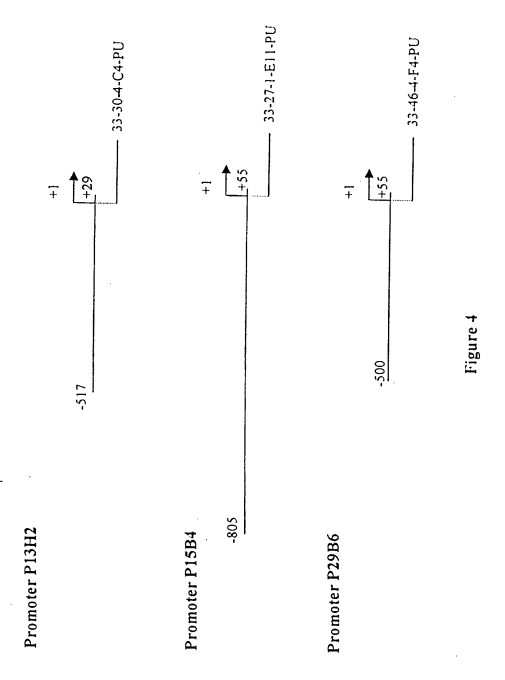


Figure 3



SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT:
 - (A) NAME : GENSET SA
 - (B) STREET : 24, RUE ROYALE
 - (C) CITY: PARIS
 - (E) COUNTRY : FRANCE
 - (F) POSTAL CODE (ZIP): 75008
- (ii) TITLE OF INVENTION: 5' ESTS FOR SECRETED PROTEINS EXPRESSED IN PROSTATE
 - (iii) NUMBER OF SEQUENCES: 593
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy Disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: Win95
 - (D) SOFTWARE: Word
- (2) INFORMATION FOR SEQ ID NO: 1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 47 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: SINGLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: Other nucleic acid
 - (ix) FEATURE:
 - (A) NAME/KEY: Cap
 - (3) LOCATION: 1
 - (D) OTHER INFORMATION: m7Gppp added to 1
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

GGCAUCCUAC UCCCAUCCAA UUCCACCCUA ACUCCUCCCA UCUCCAC

- (2) INFORMATION FOR SEQ ID NO: 2:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 46 base pairs
 - (3) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: SINGLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: Other nucleic acid
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

47

WO 99/06550 PCT/IB98/01232

(2)) INFORMATION FOR SEQ ID NO: 3:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: SINGLE (D) TOPOLOGY: LINEAR	
	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:	
ATC	CAAGAATT CGCACGAGAC CATTA	25
(2)	INFORMATION FOR SEQ ID NO: 4:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: SINGLE (D) TOPOLOGY: LINEAR	
	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:	
TAA	TGGTCTC GTGCGAATTC TTGAT	25
(2)	INFORMATION FOR SEQ ID NO: 5:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: NUCLEIC ACID - (C) STRANDEDNESS: SINGLE (D) TOPOLOGY: LINEAR	
	(ii) MOLECULE TYPE: Other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:	
CCGI	ACAAGAC CAACGTCAAG GCCGC	25
(2)	INFORMATION FOR SEQ ID NO: 6:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: SINGLE (D) TOPOLOGY: LINEAR	

WO 99/06550		3	PCT/IB98/01232
(ii) MOLEC	ULE TYPE: Other nucle	eic acid	
(xi) SEQUE	NCE DESCRIPTION: SEQ	ID NO: 6:	
TCACCAGCAG GCAGT	GGCTT AGGAG		25
(2) INFORMATION	FOR SEQ ID NO: 7:	•	
(A) (B) (C) (C)	CE CHARACTERISTICS: LENGTH: 25 base pairs TYPE: NUCLEIC ACID STRANDEDNESS: SINGLE TOPOLOGY: LINEAR		
(ii) MOLEC	ULE TYPE: Other nucle	ic acid	
(xi) SEQUE	NCE DESCRIPTION: SEQ	ID NO: 7:	
AGTGATTCCT GCTAC	TTTGG ATGGC		25
(2) INFORMATION	FOR SEQ ID NO: 8:	·	
(A) (B) (C)	CE CHARACTERISTICS: LENGTH: 25 base pairs TYPE: NUCLEIC ACID STRANDEDNESS: SINGLE TOPOLOGY: LINEAR		
(ii) MOLEC	ULE TYPE: Other nucle	ric acid	
(xi) SEQUE	NCE DESCRIPTION: SEQ	ID NO: 8:	
GCTTGGTCTT GTTCT	GGAGT TTAGA		25
(2) INFORMATION	FOR SEQ ID NO: 9:		
(A) (B) (C)	CE CHARACTERISTICS: LENGTH: 25 base pairs TYPE: NUCLEIC ACID STRANDEDNESS: SINGLE TOPOLOGY: LINEAR		
(ii) MOLEC	ULE TYPE: Other nucle	eic acid	
(xi) SEOUE	NCE DESCRIPTION: SEO	ID NO: 9:	

TCCAGAATGG GAGACAAGCC AATTT

25

(2) INFORMATION FOR SEQ ID NO: 10:

 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: SINGLE (D) TOPOLOGY: LINEAR 	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:	
AGGGAGGAGG AAACAGCGTG AGTCC	25
(2) INFORMATION FOR SEQ ID NO: 11:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 25 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: SINGLE(D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:	
ATGGGAAAGG AAAAGACTCA TATCA	25
(2) INFORMATION FOR SEQ ID NO: 12:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: SINGLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:	
AGCAGCAACA ATCAGGACAG CACAG	25
(2) INFORMATION FOR SEQ ID NO: 13:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: SINGLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:	

ATCAAGAATT CGCACGAGAC CATTA	25
(2) INFORMATION FOR SEQ ID NO: 14:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 67 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: SINGLE (D) TOPOLOGY: LINEAR 	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:	
ATCGTTGAGA CTCGTACCAG CAGAGTCACG AGAGAGACTA CACGGTACTG GTTTTTTTT	60
TTTTTVN	67
(2) INFORMATION FOR SEQ ID NO: 15:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 29 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: SINGLE(D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:	
CCAGCAGAGT CACGAGAGAG ACTACACGG	29
(2) INFORMATION FOR SEQ ID NO: 16:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: SINGLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:	
CACGAGAGAG ACTACACGGT ACTGG	25

(2) INFORMATION FOR SEQ ID NO: 17:

WO 99/06550 6 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 526 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Lymph ganglia (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(261..376) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 96 region 166..281 id N70479 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(380..486) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 54..160 id N70479 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(110..145) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 94 region 403..438 id N70479 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(196..229) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 94 region 315..348 id N70479 est (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 90..140

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 8.2

seq LLLITAILAVAVG/FP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:

AATATRARAC AGCTACAATA TTCCAGGGCC ARTCACTTGC CATTTCTCAT AACAGCGTCA 60 GAGAGAAAGA ACTGACTGAR ACGTTTGAG ATG AAG AAA GTT CTC CTC CTG ATC 113

ACA GCC ATC TTG GCA GTG GCT GTW GGT TTC CCA GTC TCT CAA GAC CAG 161 Thr Ala Ile Leu Ala Val Ala Val Gly Phe Pro Val Ser Gln Asp Gln GAA CGA GAA AAA AGA AGT ATC AGT GAC AGC GAT GAA TTA GCT TCA GGR 209 Glu Arg Glu Lys Arg Ser Ile Ser Asp Ser Asp Glu Leu Ala Ser Gly 15 WTT TTT GTG TTC CCT TAC CCA TAT CCA TTT CGC CCA CTT CCA CCA ATT 257 Xaa Phe Val Phe Pro Tyr Pro Tyr Pro Phe Arg Pro Leu Pro Pro Ile CCA TTT CCA AGA TTT CCA TGG TTT AGA CGT AAN TTT CCT ATT CCA ATA 305 Pro Phe Pro Arg Phe Pro Trp Phe Arg Arg Xaa Phe Pro Ile Pro Ile 50 CCT GAA TCT GCC CCT ACA ACT CCC CTT CCT AGC GAA AAG TAAACAARAA 354 Pro Glu Ser Ala Pro Thr Thr Pro Leu Pro Ser Glu Lys GGAAAAGTCA CRATAAACCT GGTCACCTGA AATTGAAATT GAGCCACTTC CTTGAARAAT 414 CAAAATTCCT GTTAATAAAA RAAAAACAAA TGTAATTGAA ATAGCACACA GCATTCTCTA 474 GTCAATATCT TTAGTGATCT TCTTTAATAA ACATGAAAGC AAAAAAAAA AA 526

(2) INFORMATION FOR SEQ ID NO: 18:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 17 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 1..17
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.2

seq LLLITAILAVAVG/FP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:

Met Lys Lys Val Leu Leu Ile Thr Ala Ile Leu Ala Val Ala Val 1 5 10 15

Gly

(2) INFORMATION FOR SEQ ID NO: 19:

```
(i) SEQUENCE CHARACTERISTICS:
```

- (A) LENGTH: 822 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (D) DEVELOPMENTAL STAGE: Fetal
- (F) TISSUE TYPE: kidney

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 260..464
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 153..357

id H57434

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 118..184
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 98..164

id H57434

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 56..113
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 35..92

id H57434

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 454..485
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 348..379

id H57434

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 118..545
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 1..428

id N27248

est

(ix) FEATURE:

(A) NAME/KEY: other

WO 99/06550 PCT/IB98/01232

(B) LOCATION: 65..369

(C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 41..345 id H94779 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 61..399 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 99 region 6..344 id H09880 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 408..458 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 92 region 355..405 id H09880 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 60..399 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 56..395 id H29351 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 393..432 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 90 region 391..430 id H29351 est (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 346..408 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.5 seq SFLPSALVIWTSA/AF (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19: ACTCCTTTTA GCATAGGGGC TTCGGCGCCA GCGGCCAGCG CTAGTCGGTC TGGTAAGTGC 60 CTGATGCCGA GTTCCGTCTC TCGCGTCTTT TCCTGGTCCC AGGCAAAGCG GASGNAGATC 120 CTCAAACGGC CTAGTGCTTC GCGCTTCCGG AGAAAATCAG CGGTCTAATT AATTCCTCTG 180 GTTTGTTGAA GCAGTTACCA AGAATCTTCA ACCCTTTCCC ACAAAAGCTA ATTGAGTACA 240

10	01.1270.0140.
CGTTCCTGTT GAGTACACGT TCCTGTTGAT TTACAAAAGG TGCAGGTATG AGCAGGTCT	G 300
AAGACTAACA TTTTGTGAAG TTGTAAAACA GAAAACCTGT TAGAA ATG TGG TGG TT Met Trp Trp Ph -20	
CAG CAA GGC CTC AGT TTC CTT CCT TCA GCC CTT GTA ATT TGG ACA TCT Gln Gln Gly Leu Ser Phe Leu Pro Ser Ala Leu Val Ile Trp Thr Ser -15 -5	405
GCT GCT TTC ATA TTT TCA TAC ATT ACT GCA GTA ACA CTC CAC CAT ATA Ala Ala Phe Ile Phe Ser Tyr Ile Thr Ala Val Thr Leu His His Ile 1 5 10 15	453
GAC CCG GCT TTA CCT TAT ATC AGT GAC ACT GGT ACA GTA GCT CCA RAA Asp Pro Ala Leu Pro Tyr Ile Ser Asp Thr Gly Thr Val Ala Pro Xaa 20 25 30	501
AAA TGC TTA TTT GGG GCA ATG CTA AAT ATT GCG GCA GTT TTA TGT CAA Lys Cys Leu Phe Gly Ala Met Leu Asn Ile Ala Ala Val Leu Cys Gln 35 40 45	549
AAA TAGAAATCAG GAARATAATT CAACTTAAAG AAKTTCATTT CATGACCAAA Lys	602
CTCTTCARAA ACATGTCTTT ACAAGCATAT CTCTTGTATT GCTTTCTACA CTGTTGAATT	r 662
STCTGGCAAT ATTTCTGCAG TGGAAAATTT GATTTARMTA GTTCTTGACT GATAAATATO	G 722
GTAAGGTGGG CTTTTCCCCC TGTGTAATTG GCTACTATGT CTTACTGAGC CAAGTTGTAV	v 782
TTTGAAATAA AATGATATGA GAGTGACACA AAAAAAAAAA	822

(2) INFORMATION FOR SEQ ID NO: 20:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 21 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 1..21
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.5

seq SFLPSALVIWTSA/AF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 20:

Met Trp Trp Phe Gln Gln Gly Leu Ser Phe Leu Pro Ser Ala Leu Val

WO 99/06550 PCT/IB98/01232

Ile Trp Thr Ser Ala 20

ТТТСТААААА САААААААА А

(2) INFORMATION FOR SEQ ID NO: 21:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 405 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: DOUBLE(D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Testis</pre>	
(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(103398) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 96 region 1296 id AA442893 est	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 185295 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.9</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 21:	
ATCACCTTCT TCTCCATCCT TSTCTGGGCC AGTCCCCARC CCAGTCCCTC TCCTGACCTG	60
CCCAGCCCAA GTCAGCCTTC AGCACGCGCT TTTCTGCACA CAGATATTCC AGGCCTACCT 1	.20
GGCATTCCAG GACCTCCGMA ATGATGCTCC AGTCCCTTAC AAGCGCTTCC TGGATGAGGG 1	.80
TGGC ATG GTG CTG ACC ACC CTC CCC TTG CCC TCT GCC AAC AGC CCT GTG Met Val Leu Thr Thr Leu Pro Leu Pro Ser Ala Asn Ser Pro Val -35 -30 -25	229
AAC ATG CCC ACC ACT GGC CCC AAC AGC CTG AGT TAT GCT AGC TCT GCC Asn Met Pro Thr Thr Gly Pro Asn Ser Leu Ser Tyr Ala Ser Ser Ala -20 -15 -10	277
CTG TCC CCC TGT CTG ACC GCT CCA AAK TCC CCC CGG CTT GCT ATG ATG Leu Ser Pro Cys Leu Thr Ala Pro Xaa Ser Pro Arg Leu Ala Met Met -5 1 5 10	325
CCT GAC AAC TAAATATCCT TATCCAAATC AATAAARWRA RAATCCTCCC TCCARAAGGG 3	384

405

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(2) INFORMATION FOR SEQ ID NO: 22:
```

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 37 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: $1..\overline{37}$
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq LSYASSALSPCLT/AP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 22:

Met Val Leu Thr Thr Leu Pro Leu Pro Ser Ala Asn Ser Pro Val Asn 1 5 10 15

Met Pro Thr Thr Gly Pro Asn Ser Leu Ser Tyr Ala Ser Ser Ala Leu
20 25 30

Ser Pro Cys Leu Thr 35

- (2) INFORMATION FOR SEQ ID NO: 23:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 496 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 149..331
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 1..183 id AA397994

est

- (ix) FEATURE:
 - (A) NAME/KEY: other

WO 99/06550		13	•	PCT/IB98/01
(C)	LOCATION: 328485 IDENTIFICATION METHO OTHER INFORMATION:	DD: blastn identity 96 region 179 id AA397994 est	336	
(B) (C)	URE: NAME/KEY: other LOCATION: complement IDENTIFICATION METHO OTHER INFORMATION:		28	
(B) (C)	URE: NAME/KEY: sig_peptic LOCATION: 196240 IDENTIFICATION METHO OTHER INFORMATION:			
(xi) SEQU	ENCE DESCRIPTION: SEC	Q ID NO: 23:		
AAAAAATTGG TCCC	AGTTTT CACCCTGCCG CAG	GGGCTGGC TGGG	GAGGGC AGCGGTT	TAG 60
ATTAGCCGTG GCCT	AGGCCG TTTAACGGGG TGA	ACACGAGC NTGC	AGGGCC GAGTCC#	AAGG 120
CCCGGAGATA GGAC	CAACCG TCAGGAATGC GAG	GGAATGTT TTTC	TTCGGA CTCTATO	GAG 180
GCACACAGAC AGAC	C ATG GGG ATT CTG TC Met Gly Ile Leu Ser -15			
	GAC GGC TGC AGA AAT Asp Gly Cys Arg Asn 5	Gly Ile Ala		

GAG AAG CAC AGA CTC GAG AAA TGT AGG GAA CTC GAG ASC ASC CAC TCG 327 Glu Lys His Arg Leu Glu Lys Cys Arg Glu Leu Glu Xaa Xaa His Ser 15 20 GCC CCA GGA TCA ACC CAS CAC CGA AGA AAA ACA ACC AGA AGA AAT TAT 375 Ala Pro Gly Ser Thr Xaa His Arg Arg Lys Thr Thr Arg Arg Asn Tyr 40 45

TCT TCA GCC TGAAATGAAK CCGGGATCAA ATGGTTGCTG ATCARAGCCC ATATTTAAAT 434 Ser Ser Ala

TGGAAAAGTC AAATTGASCA TTATTAAATA AAGCTTGTTT AATATGTCTC AAACAAAAAA 494 AΑ

496

(2) INFORMATION FOR SEQ ID NO: 24:

(i) SEQUENCE CHARACTERISTICS:

WO 99/06550 PCT/IB98/01232

(A) LENGTH: 15 amino acids (B) TYPE: AMINO ACID (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: PROTEIN	
(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens	
 (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 115 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.5 seq ILSTVTALTFAXA/LD 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 24:	
Met Gly Ile Leu Ser Thr Val Thr Ala Leu Thr Phe Ala Xaa Ala 1 5 10 15	
(2) INFORMATION FOR SEQ ID NO: 25: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 623 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Testis</pre>	
 (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 4996 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 10.1 seq LVLTLCTLPLAVA/SA 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 25:	
AAAGATCCCT GCAGCCCGGC AGGAGAGAAG GCTGAGCCTT CTGGCGTC ATG GAG AGG Met Glu Arg -15	57
CTC GTC CTA ACC CTG TGC ACC CTC CCG CTG GCT GTG GCG TCT GCT GGC Leu Val Leu Thr Leu Cys Thr Leu Pro Leu Ala Val Ala Ser Ala Gly -10 -5	105
TGC GCC ACG ACG CCA GCT CGC AAC CTG AGC TGC TAC CAG TGC TTC AAG Cys Ala Thr Thr Pro Ala Arg Asn Leu Ser Cys Tyr Gln Cys Phe Lys 5	153
GTC AGC AGC TGG ACG GAG TGC CCG CCC ACC TGG TGC AGC CCG CTG GAC	201

										,						
Val 20	Ser	Ser	Trp	Thr	Glu 25	Cys	Pro	Pro	Thr	Trp 30	Cys	Ser	Pro	Leu	Asp 35	
CAA Gln	GTC Val	TGC Cys	ATC Ile	TCC Ser 40	AAC Asn	GAG Glu	GTG Val	GTC Val	GTC Val 45	TCT Ser	TTT Phe	AAA Lys	TGG Trp	AGT Ser 50	GTA Val	249
														GAC Asp		297
														ATC Ile		345
														CCA Pro		393
														CCT Pro		441
														CTC Leu 130		489
					CCC Pro											534
TAAC	CACTO	TG C	GTGC	cccc	A CC	TGTG	CATI	GGC	SACCA	ACRA	CTTC	CACCO	TC I	TGGA	RACAA	594
TAAACTCTCA TGCCCCCAAA AAAAAAAA										623						

(2) INFORMATION FOR SEQ ID NO: 26:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 16 amino acids
- (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 1..16
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.1

seq LVLTLCTLPLAVA/SA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 26:

Met Glu Arg Leu Val Leu Thr Leu Cys Thr Leu Pro Leu Ala Val Ala 1 5 10 15

(2)	INFORMATION	FOR	SEO	מד	NO.	27.
. – ,	1111 0111 111 1011	- 011	JUV	10	wo.	~ / / :

(i)	SEQUENCE	CHARACTERISTICS:

- (A) LENGTH: 848 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (D) DEVELOPMENTAL STAGE: Fetal
- (F) TISSUE TYPE: kidney

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
 (B) LOCATION: 32..73
- (C) IDENTIFICATION METHOD: Von Heijne matri \mathbf{x}
- (D) OTHER INFORMATION: score 10.7

seq LWLLFFLVTAIHA/EL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:

AAC	TTTG	CCT '	TGTG	TTTT	CC A	CCCT	GAAA							TTT Phe			55
GTG Val	ACT Thr -5	GCC Ala	ATT Ile	CAT His	GCT Ala	GAA Glu 1	CTC Leu	TGT Cys	CAA Gln	CCA Pro 5	GGT Gly	GCA Ala	GAA Glu	AAT Asn	GCT Ala	r a	103
TTT Phe	AAA Lys	GTG Val	AGA Arg	CTT Leu 15	AGT Ser	ATC Ile	AGA Arg	ACA Thr	GCT Ala 20	CTG Leu	GGA Gly	GAT Asp	AAA Lys	GCA Ala 25	TAT	r :	151
GCC Ala	TGG Trp	GAT Asp	ACC Thr 30	AAT Asn	GAA Glu	GAA Glu	TAC Tyr	CTC Leu 35	TTC Phe	AAA Lys	GCG Ala	ATG Met	GTA Val 40	GCT Ala	TT(Phe	:	199
TCC Ser	ATG Met	AGA Arg 45	AAA Lys	GTT Val	CCC Pro	AAC Asn	AGA Arg 50	GAA Glu	GCA Ala	ACA Thr	GAA Glu	ATT Ile 55	TCC Ser	CAT His	GT(Val	:	247
CTA Leu	CTT Leu 60	TGC Cys	AAT Asn	GTA Val	ACC Thr	CAG Gln 65	AGG Arg	GTA Val	TCA Ser	TTC Phe	TGG Trp 70	TTT Phe	GTG Val	GTT Val	ACA Thr	<u>.</u>	295
GAC Asp 75	CCT Pro	TCA Ser	AAA Lys	AAT Asn	CAC His 80	ACC Thr	CTT Leu	CCT Pro	GCT Ala	GTT Val 85	GAG Glu	GTG Val	CAA Gln	TCA Ser	GCC Ala	;	343
ATA Ile	AGA Arg	ATG Met	AAC Asn	AAG Lys 95	AAC Asn	CGG Arg	ATC Ile	AAC Asn	AAT Asn 100	GCC Ala	TTC Phe	TTT Phe	CTA Leu	AAT Asn 105	GAC Asp	;)	391
CAA	ACT	CTG	GAA	TTT	TTA	AAA	ATC	ССТ	TCC	ACA	СТТ	GCA	CCA	CCC	ATG	;	439

									17							
Gln	Thr	Leu	Glu 110	Phe	Leu	Lys	Ile	Pro 115	Ser	Thr	Leu	Ala	Pro 120	Pro	Met	
					ATC Ile											487
					ATT Ile											535
					AAA Lys 160											583
					ATC Ile											631
					GGG Gly											679
			ACC Thr		CTC Leu	TGA	AGGGG	CTG 1	TGT	CTG	CT TO	CTC	ARA <i>I</i>	A		7 27
ATT	\AAC#	ATT T	rgtti	CTGI	G TO	SACTO	GCTG#	A GC	ATCCI	GAA	ATAC	CAAC	GAG (CAGAT	CATAT	787
WTTI	TGTI	TTC A	ACCAT	TCTI	C TI	TTGT	TAAT	AA A	TTT	SAAT	GTG	T T G#	AAA A	\AAA/	AAAAA	847
С																848

(2) INFORMATION FOR SEQ ID NO: 28:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: $1..\overline{14}$
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.7

seq LWLLFFLVTAIHA/EL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:

Met Leu Trp Leu Leu Phe Phe Leu Val Thr Ala Ile His Ala 1 5 10

- (2) INFORMATION FOR SEQ ID NO: 29:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 25 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: SINGLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: Other nucleic acid
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29:

GGGAAGATGG AGATAGTATT GCCTG

25

- (2) INFORMATION FOR SEQ ID NO: 30:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 26 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: SINGLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: Other nucleic acid
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

CTGCCATGTA CATGATAGAG AGATTC

26

- (2) INFORMATION FOR SEQ ID NO: 31:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 546 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: Genomic DNA
 - (ix) FEATURE:
 - (A) NAME/KEY: promoter
 - (B) LOCATION: 1..517
 - (ix) FEATURE:
 - (A) NAME/KEY: transcription start site
 - (B) LOCATION: 518
 - (ix) FEATURE:
 - (A) NAME/KEY: TF binding-site
 - (B) LOCATION: 17..25
 - (C) IDENTIFICATION METHOD: matinspector prediction
 - (D) OTHER INFORMATION: name CMYB 01 score 0.983 sequence TGTCAGTTG

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: complement(18..27)

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name MYOD_Q6 score 0.961

sequence CCCAACTGAC

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: complement (75..85)

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name S8_01 score 0.960

sequence AATAGAATTAG

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: 94..104

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name S8_01 score 0.966

sequence AACTAAATTAG

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: complement(129..139)

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name DELTAEF1_01 score 0.960

sequence GCACACCTCAG

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: complement(155..165)

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name GATA_C score 0.964 sequence AGATAAATCCA

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: 170..178

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name CMYB_01
score 0.958
sequence CTTCAGTTG

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: 176..189

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name GATA1_02 score 0.959

sequence TTGTAGATAGGACA

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(3) LOCATION: 180..190

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name GATA C

score 0.953 sequence AGATAGGACAT

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: 284..299
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name TAL1ALPHAE47 01

score 0.973

sequence CATAACAGATGGTAAG

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: 284..299
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name TAL1BETAE47 01

score 0.983

sequence CATAACAGATGGTAAG

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: 284..299
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name TAL1BETAITF2_01 score 0.978

sequence CATAACAGATGGTAAG

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: complement (287..296)
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name MYOD Q6 score 0.954

sequence ACCATCTGTT

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: complement (302..314)
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name GATA1 04 score 0.953

sequence TCAAGATAAAGTA

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: 393..405
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name IK1 01 score 0.963

sequence AGTTGGGAATTCC

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: 393..404
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name IK2 01 score 0.985

sequence AGTTGGGAATTC

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

WO 99/06550 PCT/IB98/01232

	(B) LOCATION: 396405 (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name CREL 01	
	score 0.962 sequence TGGGAATTCC	
(ix)	FEATURE: (A) NAME/KEY: TF binding-site (B) LOCATION: 423436 (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name GATA1_02 score 0.950 sequence TCAGTGATATGGCA	
(ix)	FEATURE: (A) NAME/KEY: TF binding-site (B) LOCATION: complement (478489) (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name SRY 02 score 0.951 sequence TAAAACAAAACA	
(ix)	FEATURE: (A) NAME/KEY: TF binding-site (B) LOCATION: 486493 (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name E2F_02 score 0.957 sequence TTTAGCGC	
(ix)	FEATURE: (A) NAME/KEY: TF binding-site (B) LOCATION: complement(514521) (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name MZF1_01 score 0.975 sequence TGAGGGGA	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 31:	
TGAGTGGAGT	GTTACATGTC AGTTGGGTTA AGTTTGTTAA TGTCATTCAA ATCTTCTATG	60
TCTTGATTTG	CCTGCTAATT CTATTATTTC TGGAACTAAA TTAGTTTGAT GGTTCTATTA	120
GTTATTGACT	GAGGTGTGCT AATCTCCCAT TATGTGGATT TATCTATTTC TTCAGTTGTA	180
GATAGGACAT	TGATAGATAC ATAAGTACCA GGACAAAAGC AGGGAGATCT TTTTTCCAAA	240
ATCAGGAGAA	AAAAATGACA TCTGGAAAAC CTATAGGGAA AGGCATAACA GATGGTAAGG	300
ATACTTTATC	TTGAGTAGGA GAGCCTTCCT GTGGCAACGT GGAGAAGGGA AGAGGTCGTA	360
GAATTGAGGA	GTCAGCTCAG TTAGAAGCAG GGAGTTGGGA ATTCCGTTCA TGTGATTTAG	420
CATCAGTGAT	ATGGCAAATG TGGGACTAAG GGTAGTGATC AGAGGGTTAA AATTGTGTGT	480
TTTGTTTTAG	CGCTGCTGGG GCATCGCCTT GGGTCCCCTC AAACAGATTC CCATGAATCT	540
CTTCAT		546

22

- (2) INFORMATION FOR SEQ ID NO: 32:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: SINGLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: Other nucleic acid
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 32:

GTACCAGGGA CTGTGACCAT TGC

23

- (2) INFORMATION FOR SEQ ID NO: 33:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 24 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: SINGLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: Other nucleic acid
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 33:

CTGTGACCAT TGCTCCCAAG AGAG

24

- (2) INFORMATION FOR SEQ ID NO: 34:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 861 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: Genomic DNA
 - (ix) FEATURE:
 - (A) NAME/KEY: promoter
 - (B) LOCATION: 1..806
 - (ix) FEATURE:
 - (A) NAME/KEY: transcription start site
 - (B) LOCATION: 807
 - (ix) FEATURE:
 - (A) NAME/KEY: TF binding-site
 - (B) LOCATION: complement(60..70)
 - (C) IDENTIFICATION METHOD: matinspector prediction
 - (D) OTHER INFORMATION: name NFY_Q6
 score 0.956
 sequence GGACCAATCAT

23 (ix) FEATURE: (A) NAME/KEY: TF binding-site (B) LOCATION: 70..77 (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name MZF1 01 score 0.962 sequence CCTGGGGA (ix) FEATURE: (A) NAME/KEY: TF binding-site (B) LOCATION: 124..132 (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name CMYB 01 score $0.9\overline{9}4$ sequence TGACCGTTG (ix) FEATURE: (A) NAME/KEY: TF binding-site (B) LOCATION: complement(126..134) (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name VMYB 02 score $0.9\overline{8}5$ sequence TCCAACGGT (ix) FEATURE: (A) NAME/KEY: TF binding-site (B) LOCATION: 135..143 (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name STAT 01 score $0.9\overline{68}$ sequence TTCCTGGAA (ix) FEATURE: (A) NAME/KEY: TF binding-site (B) LOCATION: complement(135..143) (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name STAT 01 score $0.9\overline{5}1$ sequence TTCCAGGAA (ix) FEATURE: (A) NAME/KEY: TF binding-site (B) LOCATION: complement (252..259) (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name MZF1 01 score $0.9\overline{5}6$ sequence TTGGGGGA (ix) FEATURE: (A) NAME/KEY: TF binding-site (B) LOCATION: 357..368 (C) IDENTIFICATION METHOD: matinspector prediction (D) OTHER INFORMATION: name IK2 01 score 0.965 sequence GAATGGGATTTC

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: 384..391
- (C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name MZF1_01 score 0.986

sequence AGAGGGGA

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: complement (410..421)

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name SRY 02

score 0.955

sequence GAAAACAAAACA

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: 592..599

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name MZF1_01

score 0.960

sequence GAAGGGGA

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: 618..627

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name MYOD_Q6

score 0.981

sequence AGCATCTGCC

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: 632..642

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name DELTAEF1 01

score 0.958

sequence TCCCACCTTCC

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: complement(813..823)

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name S8 01

score 0.992

sequence GAGGCAATTAT

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: complement (824..831)

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name MZF1 01

score 0.986

sequence AGAGGGGA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 34:

TACTATAGGG CACGCGTGGT CGACGGCCGG GCTGTTCTGG AGCAGAGGGC ATGTCAGTAA 60

TGATTGGTCC CTGGGGAAGG TCTGGCTGGC TCCAGCACAG TGAGGCATTT AGGTATCTCT 120

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CTCAGAGGGC	TAGGCACGAG	GGAAGGTCAG	AGGAGAAGGS	AGGSARGGCC	CAGTGAGARG	240
GGAGCATGCC	TTCCCCCAAC	CCTGGCTTSC	YCTTGGYMAM	AGGGCGKTTY	TGGGMACTTR	300
AAYTCAGGGC	CCAASCAGAA	SCACAGGCCC	AKTCNTGGCT	SMAAGCACAA	TAGCCTGAAT	360
GGGATTTCAG	GTTAGNCAGG	GTGAGAGGGG	AGGCTCTCTG	GCTTAGTTTT	GTTTTGTTTT	420
CCAAATCAAG	GTAACTTGCT	CCCTTCTGCT	ACGGGCCTTG	GTCTTGGCTT	GTCCTCACCC	480
AGTCGGAACT	CCCTACCACT	TTCAGGAGAG	TGGTTTTAGG	CCCGTGGGGC	TGTTCTGTTC	540
CAAGCAGTGT	GAGAACATGG	CTGGTAGAGG	CTCTAGCTGT	GTGCGGGGCC	TGAAGGGGAG	600
TGGGTTCTCG	CCCAAAGAGC	ATCTGCCCAT	TTCCCACCTT	CCCTTCTCCC	ACCAGAAGCT	660
TGCCTGAGCT	GTTTGGACAA	AAATCCAAAC	CCCACTTGGC	TACTCTGGCC	TGGCTTCAGC	720
TTGGAACCCA	ATACCTAGGC	TTACAGGCCA	TCCTGAGCCA	GGGGCCTCTG	GAAATTCTCT	780
TCCTGATGGT	CCTTTAGGTT	TGGGCACAAA	ATATAATTGC	СТСТССССТС	TCCCATTTTC	840
TCTCTTGGGA	GCAATGGTCA	С				861

(2) INFORMATION FOR SEQ ID NO: 35:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: SINGLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 35:

CTGGGATGGA AGGCACGGTA

20

(2) INFORMATION FOR SEQ ID NO: 36:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20 base pairs

 - (B) TYPE: NUCLEIC ACID
 (C) STRANDEDNESS: SINGLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 36:

GAGACCACAC AGCTAGACAA

20

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(i) SEQUENCE CHARACTERISTICS:
```

- (A) LENGTH: 555 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: Genomic DNA

(ix) FEATURE:

- (A) NAME/KEY: promoter
- (B) LOCATION: 1..500

(ix) FEATURE:

- (A) NAME/KEY: transcription start site
- (B) LOCATION: 501

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: 191..206
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name ARNT 01

score 0.964

sequence GGACTCACGTGCTGCT

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: 193..204
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name NMYC_01

score 0.965

sequence ACTCACGTGCTG

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: 193..204
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name USF_01

score 0.985

sequence ACTCACGTGCTG

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: complement(193..204)
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name USF 01

score $0.\overline{9}85$

sequence CAGCACGTGAGT

(ix) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: complement(193..204)
- (C) IDENTIFICATION METHOD: matinspector prediction
- (D) OTHER INFORMATION: name NMYC_01

score 0.956

sequence CAGCACGTGAGT

(iz) FEATURE:

- (A) NAME/KEY: TF binding-site
- (B) LOCATION: complement (193..204)
- (C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name MYCMAX 02

score 0.972

sequence CAGCACGTGAGT

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: 195..202

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name USF_C

score 0.997

sequence TCACGTGC

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: complement(195..202)

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name USF_C score 0.991

sequence GCACGTGA

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: complement(210..217)

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name MZF1_01 score 0.968

sequence CATGGGGA

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: 397..410

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name ELK1_02 score 0.963

sequence CTCTCCGGAAGCCT

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: 400..409

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name CETS1P54_01

score 0.974

sequence TCCGGAAGCC

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: complement(460..470)

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name AP1_Q4

score $0.\overline{9}63$

sequence AGTGACTGAAC

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: complement (460..470)

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name APIFJ 02

score 0.961

sequence AGTGACTGAAC

(ix) FEATURE:

(A) NAME/KEY: TF binding-site

(B) LOCATION: 547..555

(C) IDENTIFICATION METHOD: matinspector prediction

(D) OTHER INFORMATION: name PADS_C score 1.000 sequence TGTGGTCTC

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 37:

CTATAGGGCA CGCKTGGTCG ACGGCCCGGG CTGGTCTGGT CTGTKGTGGA GTCGGGTTGA 60 AGGACAGCAT TTGTKACATC TGGTCTACTG CACCTTCCCT CTGCCGTGCA CTTGGCCTTT 120 KAWAAGCTCA GCACCGGTGC CCATCACAGG GCCGGCAGCA CACACATCCC ATTACTCAGA 180 AGGAACTGAC GGACTCACGT GCTGCTCCGT CCCCATGAGC TCAGTGGACC TGTCTATGTA GAGCAGTCAG ACAGTGCCTG GGATAGAGTG AGAGTTCAGC CAGTAAATCC AAGTGATTGT CATTCCTGTC TGCATTAGTA ACTCCCAACC TAGATGTGAA AACTTAGTTC TTTCTCATAG 360 GTTGCTCTGC CCATGGTCCC ACTGCAGACC CAGGCACTCT CCGGAAGCCT GGAAATCACC 420 CGTGTCTTCT GCCTGCTCCC GCTCACATCC CACACTTGTG TTCAGTCACT GAGTTACAGA 480 TTTTGCCTCC TCAATTTCTC TTGTCTTAGT CCCATCCTCT GTTCCCCTGG CCAGTTTGTC 540 TAGCTGTGTG GTCTC 555

(2) INFORMATION FOR SEQ ID NO: 38:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 120 base pairs
 - (3) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 16..84
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 11.4

seq VLALLLFVHYSNG/DE

(xi) SEQUENCE DESCRIPTION: SEQ ID MO: 33:

ACTICCTGGT GCTGC ATG GTG TTC-GTG CAC CTG TAC CTG GGT AAC GTG CTG

Met Val Phe Val His Leu Tyr Leu Gly Asn Val Leu

-20 -15

GGG GTG GTG GTG GTG GAC TAC AGC AAC GGC GAC GAA AGC AGC GAT 99

WO 77/00330		29	PC1/1B98/
Ala Leu Leu Leu -10	Phe Val His Tyr -5	Ser Asn Gly Asp Glu l	Ser Ser Asp 5
CCC GGG CCC CAR Pro Gly Pro Gln			120
	FOR SEQ ID NO: 3		
(A) (B) (C)	NCE CHARACTERISTI LENGTH: 303 base TYPE: NUCLEIC AC STRANDEDNESS: DO TOPOLOGY: LINEAR	e pairs CID OUBLE	
(ii) MOLE	CULE TYPE: CDNA		
(A)	INAL SOURCE: ORGANISM: Homo S TISSUE TYPE: Nor		
(B) (C)	NAME/KEY: sig_pe LOCATION: 2022	88 ETHOD: Von Heijne ma	
(xi) SEQUI	ENCE DESCRIPTION:	SEQ ID NO: 39:	
AAAAGTGGAA AATG	GGAGGC ATGAAATACA	TCTTTTCGTT GTTGTTCT	TT CTTTTGCTAG 60
AAGGAGGCAA AACA	GAGCAA GTAAAACATT	CAGAGACATA TTGCATGT	TT CAAGACAAGA 120
AGTACAGAGT GGGT	SAGAGA TGGCATCCTT	ACCTGGAACC TTATGGGT	TG GTTTACTGCG 180
rgaactgcat ctgc		ATG TGC TTT GCA GCC. Met Cys Phe Ala Ala -25	
GTC CAA ATG TTC Val Gln Met Phe	ATT GCC TTT CTC Ile Ala Phe Leu -15	CTG TGC ATA TTC CTC Leu Cys Ile Phe Leu -10	ATC TGT GCT 279 Ile Cys Ala -5
	CAG AAG AGT GGG Gln Lys Ser Gly 5		303
(2) INFORMATION	FOR SEQ ID NO: 4	0:	
(A) (B) (C)	NCE CHARACTERISTI LENGTH: 313 base TYPE: NUCLEIC AC STRANDEDNESS: DO TOPOLOGY: LINEAR	pairs ID UBLE	

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 203..280

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 11

seq VLFLFLFWGVSLA/GS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 40:

AAGGATGCTA TGCAAGTCAC TAATAAAGGA AGACACGGAC AGATGAACTT AAAAGAGAAG 60
CTTTAGCTGC CAAAGATTGG GAAAGGGAAA GGMCAAAAAA GACCCCTGGG CTACACGGCG 120

TAGGTGCAGG GTTTCCTACT GCTGTTCTTT TATGCTGGGA GCTGTGGCTG TAACCAACTA 180

GGAAATAACG TATGCAGCAG CT ATG GCT GTC AGA GAG TTG TGC TTC TCA AGA

Met Ala Val Arg Glu Leu Cys Phe Ser Arg

-25

-20

CAA AGG CAA GTC CTG TTT CTT TTT CTT TTT TGG GGA GTG TCC TTG GCA
Gln Arg Gln Val Leu Phe Leu Phe Leu Phe Trp Gly Val Ser Leu Ala
-15 -5

GGT TCT GGG TTT GGA CGT TAT TCG GTG ACC GGG
Gly Ser Gly Phe Gly Arg Tyr Ser Val Thr Gly

1 5 10

(2) INFORMATION FOR SEQ ID NO: 41:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 323 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 117..170
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.7

seq LILLALATGLVGG/ET

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 41:

AGAGCBNMAG CCCCAGAGCC TAGGAACCTG GGGCCCGCTC	CTCCCCCCTC CAGGCC ATG 11
AGG ATT CTG CAG TTA ATC CTG CTT GCT CTG GCA Arg Ile Leu Gln Leu Ile Leu Leu Ala Leu Ala -15	ACA GGG CTT GTA GGG 16 Thr Gly Leu Val Gly -5
GGA GAG ACC AGG ATC ATC AAG GGG TTC GAG TGC Gly Glu Thr Arg Ile Ile Lys Gly Phe Glu Cys 1 5 10	AAG CCT CAC TCC CAG 21: Lys Pro His Ser Gln 15
CCC TGG CAG GCA GCC CTG TTC GAG AAG ACG CGG Pro Trp Gln Ala Ala Leu Phe Glu Lys Thr Arg 20 25	CTA CTC TGT GGG GCG 26: Leu Leu Cys Gly Ala 30
ACG CTC ATC GCC CCC AGA TGG CTC CTG ACA GCA Thr Leu Ile Ala Pro Arg Trp Leu Leu Thr Ala 35 40	GCC CAC TGC CTC AAG 31: Ala His Cys Leu Lys 45
CCC CGC TAC GGG Pro Arg Tyr Gly 50	32:
(2) INFORMATION FOR SEQ ID NO: 42: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 264 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Hypertrophic pr (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 94147 (C) IDENTIFICATION METHOD: Von H (D) OTHER INFORMATION: score 10 seq LILL (xi) SEQUENCE DESCRIPTION: SEQ ID NO:	eijne matrix .7 ALATGLVGG/ET
AAGAGGTTGA GGTGGCTGCG GGACTGGAAG TCATCGGGCA	
AAACCTGGGG CCCGCTCCTC CCCCCTCCAG GCC ATG AGG Met Arg	G ATT CTG CAG TTA ATC 114 I lle Leu Gln Leu Ile -15
CTG CTT GCT CTG GCA ACA GGG CTT GTA GGG GGA Leu Leu Ala Leu Ala Thr Gly Leu Val Gly Gly -10	
AAG GGG TTC GAG TGC AAG CCT CAC TNC CAG CCC	TGG CAG GCA GCC CTG 210

Lys	s Gly	/ Phe	e Glu	Cys 10	Lys	Pro	His	Xaa	Gln 15	Pro	Trp	Gln	Ala	A1a 20	Leu	
TTC Phe	C GAG ∋ Glu	AAC Lys	ACG Thr 25	CGG Arg	CTA Leu	CTC Leu	TGT Cys	GGG Gly 30	Ala	ACG Thr	CTC Leu	ATC Ile	GCC Ala 35	CCC	AGA Arg	258
	CTC Leu															264
(2)	INF	ORMA	TION	FOR	SEQ	ID 1	10:	43:								
	- (i) S	(B) (C)	NCE (LENG TYPE STRA TOPO	TH: : NU NDEC	331 CLEI NESS	base C AC	e pai CID OUBLE								
	(ii) I	MOLEC	ULE	TYPE	: CE	NA									
	(,	vi) (ORIGI (A) (F)	NAL ORGA TISS	NISM	l: Ho	mo S Hyp	Sapie ertr	ns ophi	.c pr	osta	ıte				
			(B) (C)	NAME LOCA IDEN OTHE	TION TIFI R IN	: 23 CATI FORM	11 ON M ATIO	2 ETHC N:	D: V scor seq	e 10 SLLL	.6 AVLV	e ma				
CTC'	TAGA	ACC (GACC	CACC	A CC	ATG Met	AGG Arg	TCC	TGC	CTG	TGG	AGA	TGC	AGO	CAC	
						-30		001	Cys	Leu	-25		Cys	n.	J His	52
CTG Leu -20	AGC Ser	CAA Gln	GGC Gly	GTC Val	CAG Gln -15	-30 TGG	TCC	TTG	СТТ	стс	-25 GCT	GTC	C TC	CTC	ጥጥ	100
-20	CTC	TTC	GGC	GTC Val TTG	-15 CCC	-30 TGG Trp TCT	TCC Ser DNH	TTG Leu AVT	CTT Leu TRR	CTG Leu -10	-25 GCT Ala	GTC Val	CTG Leu	GTC Val	TTC Phe -5	
-20 TTT Phe	CTC Leu	TTC Phe	GGC Gly GCC	GTC Val TTG Leu 1	-15 CCC Pro	TGG Trp TCT Ser	TCC Ser DNH Xaa	TTG Leu AVT Xaa 5	CTT Leu TRR Xaa	CTG Leu -10 KGD Xaa	-25 GCT Ala SCT Xaa	GTC Val CAA Gln	CTG Leu ACA Thr 10	GTC Val AAG Lys	TTC Phe -5 CCT Pro	100
TTT Phe	CTC Leu AGG Arg	TTC Phe CAT His 15	GGC Gly GCC Ala	GTC Val TTG Leu 1 CGC	CCC Pro	TGG Trp TCT Ser:	TCC Ser DNH Xaa AAC Asn 20	TTG Leu AVT Xaa 5 ATT Ile	CTT Leu TRR Xaa AAA Lys	CTG Leu -10 KGD Xaa GAA Glu	-25 GCT Ala SCT Xaa AGG Arg	GTC Val CAA Gln TCT Ser 25	CTG Leu ACA Thr 10 CTA Leu	GTC Val AAG Lys CWG Xaa	TTC Phe -5 CCT Pro TCC Ser	100
-20 TTT Phe TCC Ser CTG Leu	CTC Leu AGG Arg GCA Ala 30	TTC Phe CAT His 15 AAG Lys	GGC Gly GCC Ala CAA Gln	GTC Val TTG Leu 1 CGC Arg	CCC Pro ACA Thr TCC Ser	TGG Trp TCT Ser GAG Glu CAG Gln 35	TCC Ser DNH Xaa AAC Asn 20 GCA Ala	TTG Leu AVT Xaa 5 ATT Ile CCC Pro	CTT Leu TRR Xaa AAA Lys ACA Thr	CTG Leu -10 KGD Xaa GAA Glu AGG Arg	-25 GCT Ala SCT Xaa AGG Arg	GTC Val CAA Gln TCT Ser 25 AGG	CTG Leu ACA Thr 10 CTA Leu	GTC Val AAG Lys CWG Xaa ACA Thr	TTC Phe -5 CCT Pro TCC Ser ACC Thr	100 148 196

Gln Pro Lys Ala His Thr Thr Gly Asp Arg Arg Lys Gly
65 70

121	INFORMATION	FOR	SEO	TD	NO.	11.
(ム)	INCOMMITON	LOK	SEQ	עב	NO:	44:

1:1	CEOHENCE	CHARACTERISTICS .
(1)	SEQUENCE.	-CHARACTERISTICS+

- (A) LENGTH: 406 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Cancerous prostate

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 167..220
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 10.6

seq XILLALATGLVGG/EI

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 44:

AATGTGGGAC GTGGC	TTTGT TCTAATAAGA CG	GAAGGGTGG AGTGCAGGCT TGGAAAGCAG	60
GAGAGCTCAG CCTAC	GTCTT TAATCCTCCT GC	CCCACCCCT TGGRTTCTGT CTCCACTGGG	120
RCTCAAGASV AGGAC	CCTGG GGGCCCGCTC CT	TCCCCCCTC CAGGCC ATG AGG ATT Met Arg Ile	175
		A ACA GGG CTT GTA GGG GGA GAG a Thr Gly Leu Val Gly Gly Glu -5 1	223
ATC AGG ATC ATC ATC Ile Arg Ile Ile 5	AAG GGG TTC GAG TGC Lys Gly Phe Glu Cys 10	C AAG CCT CAC TCC CAG CCC TGG s Lys Pro His Ser Gln Pro Trp 0 15	271
		G CTA CTA CTG TGG GGC GAC GCT g Leu Leu Trp Gly Asp Ala 30	319
		C AGC CCA CTG CCT CAA GCC CCG r Ser Pro Leu Pro Gln Ala Pro 45	367
	CCT GGG GCA GCA CAA Pro Gly Ala Ala Gln 55		406

(2) INFORMATION FOR SEQ ID NO: 45:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 187 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: DOUBLE(D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 35148 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 10.4</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 45:	
ATACTGTTTA TAAGCAACCT TGGTTTTACA TAGT ATG TTG GAA GAG TGT GGG GCT Met Leu Glu Glu Cys Gly Ala -35	55
GGG GTT GAT TTA GGA TTT GGA GGT GTA AAG TTT GCC AGT GAG ACA CCA Gly Val Asp Leu Gly Phe Gly Gly Val Lys Phe Ala Ser Glu Thr Pro -30 -25 -20	103
AAC CTT CTC TGG CTG CTT TTA AAA CTK GTA AGT ACC YCT TGG GCT GTA Asn Leu Leu Trp Leu Leu Lys Leu Val Ser Thr Xaa Trp Ala Val -10 -5 1	151
AGA GTG ACT TTG ATC ATA TTT AAC AAC CAG GCA AGG Arg Val Thr Leu Ile Ile Phe Asn Asn Gln Ala Arg 5 10	187
(2) INFORMATION FOR SEQ ID NO: 46:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 329 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 249317 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 10.2</pre>	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 46:

ATCTACTATA AAATCGATAG AAAAAAAAGT TCTTTATGGC TACTGGTCAG CTTTTATTCC	60
TGATACGCCT GAACTTGGCA GCCCACAGTC AGTGTCCTTG ATGACTCTTA SATTGAAAGA	120
CCCKTCTTCC AAAGACACGT GCCTGTGCTC TGCAAGTTTK ATCTGCCATC TTGGAAGGCT	180
CAAAGCAGTT TCTTTCTGTT GCTGAAGATA CCAGTGACCA CAGAAGGGCT TTTACCCCCT	240
TCTCCGTA ATG ATC GCT TGC AGC ATT AGA GAG TTG CAC AGA TGT CTK TTG Met Ile Ala Cys Ser Ile Arg Glu Leu His Arg Cys Leu Leu -20 -15 -10	290
TTA GCT TTG GTG GCG GAG TCA TCC TCA CAG ACC CAC GGG Leu Ala Leu Val Ala Glu Ser Ser Gln Thr His Gly -5	329
(2) INFORMATION FOR SEQ ID NO: 47:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 277 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Cancerous prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 182232 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 10.2</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 47:	
AGTTTTTTCC AGCTCCTGGG CGAATCCCAC ATCTGTTTCA ACTCTCCGCC GAGGGCGAGC	60
AGGAGCGAGA GTGTGTCGAA TCTGCGAGTG AAGAGGGAAC SAGGGGAAAA GAAACAAAGC	120
CACAGACGCA ACTTGAGACT CCCGCATCCC AAAAGAAGCA CCAGATCAGC AAAAAAAAGAA	180
G ATG GGC CCC CCG AGC CTC GTG CTG TGC TTG CTG TCC GCA ACT GTG TTC Met Gly Pro Pro Ser Leu Val Leu Cys Leu Leu Ser Ala Thr Val Phe -15 -5	229
TCC CTG CAG GGT GGA AGC TCG GCC TTC CTG TCG CAC CAC CGC CCC GGG Ser Leu Gln Gly Gly Ser Ser Ala Phe Leu Ser His His Arg Pro Gly 1 5 10	277

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 352 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Hypertrophic prostate (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 17..121 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 9 seq AMWWLLLWGVLOX/XP (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 48: AGATGTCCAG TTCCAG ATG CCT GGA CCC AGA GTG TGG GGG AAA TAT CTC TGG Met Pro Gly Pro Arg Val Trp Gly Lys Tyr Leu Trp -35 AGA AGC CCT CAC TCC AAA GGC TGT CCA GGC GCA ATG TGG TGG CTG CTT 100 Arg Ser Pro His Ser Lys Gly Cys Pro Gly Ala Met Trp Trp Leu Leu -15CTC TGG GGA GTC CTC CAG GST TKG CCC AAC CCG GGG CTC CGT CCT CTT 148 Leu Trp Gly Val Leu Gln Xaa Xaa Pro Asn Pro Gly Leu Arg Pro Leu GGC CHA AGA GCT ACC CCA GCA GCT GAC ATC CCC CGG GTA CCC AGA GCC Gly Xaa Arg Ala Thr Pro Ala Ala Asp Ile Pro Arg Val Pro Arg Ala 15 GTA TGG CAA AGG CCA AGA GAG CAR CAC GGA CAT CAA GGC TCC AGA GGG 244 Val Trp Gln Arg Pro Arg Glu Gln His Gly His Gln Gly Ser Arg Gly 30 CTT TGC TGT GAG GCT CGT CTT CCA GGA CTT CGA CCT GGA GCC GTC CCA 292 Leu Cys Cys Glu Ala Arg Leu Pro Gly Leu Arg Pro Gly Ala Val Pro GGA CTG TGC AGG GGA CTC TGT CAC AAT CTC ATT CGT CGG TTC GGA TCC 340 Gly Leu Cys Arg Gly Leu Cys His Asn Leu Ile Arg Arg Phe Gly Ser 65 AAG CCA CTC GGG 352 Lys Pro Leu Gly 75 (2) INFORMATION FOR SEQ ID NO: 49:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 450 base pairs

V	VO 99)/0655	50						37	,		٠			PCT	/ IB 98/
			(C)	STRA	ANDE	JCLEI ONESS	6: D0	OUBLE								
	(:	ii) 1	MOLE	CULE	TYP	E: CI	ANC									
	(7	zi) (ORG	ANISM	RCE: 1: Ho TYPE:				state	2					
	i)	ix) 1	(B) (C)	NAME LOCA I DEN	TION TIFI	(: si N: 15 [CAT] NFORM	512 ION N	216 4ETHC	D: V		. 8					
	()	(i) 5	SEQUE	ENCE	DESC	CRIPT	CION	: SE() ID	NO:	49:					
AAG <i>I</i>	AGCC	CCA (CGGC	CAGCT	rc c	rtcc:	rgtto	c cc	CTGG	CGGC	CCC	rcgc'	rtc '	rtcc:	TTCTGG	60
ATG	GGGG	CCC I	AGGG	GCC	AG GA	AGAG	LATA	A ASC	SSGWI	KDKG	GAR	GGT	GCC (CGGC	ACAACC	120
AGAG	CGCCC	CAG '	TCAC	AGGC	GA GA	AGCC	CTGG				g Pro				G CTG Leu -15	174
CTG Leu	CTG Leu	CTC Leu	ACG Thr	CTT Leu -10	GCC Ala	CTC Leu	CTG Leu	GGG Gly	GGC Gly -5	CCC Pro	ACC Thr	TGG Trp	GMA Xaa	SGG Xaa 1	AAG Lys	222
			CCT Pro													270
			ATC Ile													318
AAA Lys 35	AGT Ser	GTC Val	CAG Gln	GTG Val	AAA Lys 40	CTT Leu	GGA Gly	GAC Asp	TCC Ser	TGG Trp 45	GAC Asp	GTG Val	AAA Lys	CTG Leu	GGA Gly 50	366
			TGG Trp													414
			AGT Ser 70													450
(2)	INFO	ORMA'	TION	FOR	SEQ	ID 1	NO: !	50:								
	()	L) Si	EQUEN	ICE (CHARA	ACTE	RIST	ICS:								

(2) INFO

(A) LENGTH: 181 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

			•	•						38						CITIDA
	(i	i)	MOLE	CULE	TYP	E: C	DNA									
	(v	ri)	(A)	INAL ORG TIS	ANIS	M: H	omo	Sapi osta	ens te							
	(i	ж)	(B) (C)	URE: NAM LOC. IDE: OTH	ATIO NTIF	N: 5 ICAT	49 ION	METH	OD:	re 8	. 6	ne m				
	(×	i) \$	SEQU	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	50:					
AGA	C ATG Met -15	٧a.	A AG' l Se:	r GTe	G AG	T TT. r Le -1	u Al	G CT a Le	G CT u Le	G TC	C GG. r Gl -	y Tr	G GT p Va	T GG	T AGC y Ser	49
AGA Arg 1	CAG Gln	GGT Gly	GGA Gly	GTA Val 5	GGG Gly	TTA Leu	AGC Ser	ACA Thr	CTG Leu 10	GTC Val	ACC Thr	TTA Leu	GGA Gly	TTG Leu 15	GTT Val	97
TCC Ser	TGG Trp	TGC Cys	TGG Trp 20	AGA Arg	ATG Met	GTT Val	AGG Arg	ACA Thr 25	CAG Gln	GCC Ala	TTG Leu	GAA Glu	GGT Gly 30	TTT Phe	TTG Leu	145
AGT Ser	GTG A	AAA Lys 35	TAT Tyr	TAC Tyr	TCA Ser	GCG Ala	TTT Phe 40	TCT Ser	GCA Ala	GAC Asp	CTG Leu					181
(2)	INFO	RMAT	'ION	FOR	SEQ	ID N	10: 5	51:								
	(i)	SE	OUEN	CE C	HARA	CTER	ייי פיזי	.cs.								
			(A)	LENG	TH:	293	base	pai	rs							
				TYPE STRA												
	•			TOPO					•							
	(ii	.) M	OLEC	ULE	TYPE	: CD	NA									
	(vi	.) 0	RIGI	NAL	SOUR	CE:										
				ORGA			mo S	apie	ns							

- (F) TISSUE TYPE: Prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 129..275
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.5

seq IVFLLLRVSPCLG/PS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 51:

1	•
4	٠,

ATAAAGCCTT CCTTTAAAGC TTTATAATAA TCATATTTAT TAATAATGCT GTTGTGCATA	120
CTTATAGT ATG CAT ATA TTC AGC ATA TGT TGC ATG TST TCA GAA TTA CAT Met His Ile Phe Ser Ile Cys Cys Met Xaa Ser Glu Leu His -45 -40	170
AAG ATG AAA TCC CTT TCA TTG CAA CTT GCA AGT GAG AAA AGA TCC TTA Lys Met Lys Ser Leu Ser Leu Gln Leu Ala Ser Glu Lys Arg Ser Leu -35 -20	218
GTG GCT CTG GTG GAA GAA ATA GTA TTT CTT CTC AGG GTG TCT CCC Val Ala Leu Val Glu Glu Ile Val Phe Leu Leu Arg Val Ser Pro -15 -10 -5	266
TGC CTT GGC CCC TCC CAB AAG CCC CGG Cys Leu Gly Pro Ser Xaa Lys Pro Arg 1 5	293
(2) INFORMATION FOR SEQ ID NO: 52:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 323 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: DOUBLE(D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 258308 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 8.3</pre>	
(%i) SEQUENCE DESCRIPTION: SEQ ID NO: 52:	
AGCGCCGAGC TGACCGGGCG ACGCCGCGGG AGGTTCTGGA AACGCCGGGA GCTGCGAGTG	60
TCCAGACATC CTTGTGGAAC CAGGCGTTGT KTTTCCTTGG CAGCTGCGGA GACCCGTGAT	120
AATTCGTTAA CTAATTCAAC AAACGGGACC CTTCTGTGTG CCAGAAACCG CAAGCAGTTG	180
CTAACCCAGT GGGACAGGCG GATTGGAAGA GCGGGAAGGT CCTGGCCCAG AGCAGTGTGA	240
CACTTCCCTC TGTGACC ATG AAA CTC TGG GTG TCT GCA TTG CTG ATG GCC Met Lys Leu Trp Val Ser Ala Leu Leu Met Ala -15 -10	290
TGG TTT GGT GTC CTG AGC TGT GTG CAG ACC GGG Trp Phe Gly Val Leu Ser Cys Val Gln Thr Gly -5 1 5	323

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(2)	INF	ORM	OITA	N FOR	R SEQ) ID	NO:	53:								
	(:	i) :	(B) (C)	LEN TYP STR	CHAR IGTH: PE: N ANDE OLOG	235 UCLE DNES	bas IC A SS: D	e pa CID OUBL	irs							
	(i	Li)	MOLE	CULE	TYP	E: C	DNA									
	(1	/i)	ORIG (A) (F)	ORG	SOU ANISI SUE	м: н	omo	Sapi rmal	ens pro	stat	e					
	(i	.x)	(B) (C)	NAM LOC.	E/KE ATION NTIF: ER IN	N: 9 ICAT	21 ION	57 METH	0D: '	re 8	Heij .3 PLML					
	(x	:i)	SEQU	ENCE	DES	CRIP	TION	: SE	QID	NO:	53:					
AGAC	CTGA	GT	CATC	CCCA	GG G	АТСА	GGAG	С СТ	CCAG	CAGG	GAA	CCTT	CCA	TTAT	ATTCTT	60
CAAG	CAAC	TT	ACAG	CTGC.	AC CO	GACA	GTTG	C G	ATG :	Lys	GTT (Val :	CTA . Leu	ATC Ile	TCT '	TCC Ser	112
CTC Leu -15	CTC Leu	CTG Leu	TTG Leu	CTG Leu	CCA Pro -10	CTA Leu	ATG Met	CTG Leu	ATG Met	TCC Ser -5	ATG Met	GTC Val	TCT Ser	AGC Ser	AGC Ser 1	160
CTG /	AWT Xaa	CCA Pro	GGG Gly 5	GTC Val	GCC Ala	AGA Arg	GGC Gly	CAC His 10	AGG Arg	GAC Asp	CGA Arg	GGC Gly	CAG Gln 15	GCT Ala	TCT Ser	208
AGG A	AGA Arg	TGG Trp 20	CTC Leu	CAG Gln	GAA Glu	GGC Gly	GGA Gly 25	CTG Leu								235
(2)			rion													
	(i)) SI	(B) (C)	LENG TYPE STRA	HARA TH: : NU NDED LOGY	365 CLEI NESS	base C AC : DC	pai ID UBLE								
	(1)	i) 1	10LEC	ULE	TYPE	: CE	ANG									
	(v)	L) (ORIGI (A) (F)	ORGA	SOUR NISM UE T	: Но	mo S Can	apie cero	ns us p	rost	ate					

- (B) LOCATION: 159..224
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 8.3

seq LLLPLMLMSMVSS/SL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 54:

ACTGTTCTCG CCCTCAAATG GGAACGCTGA CCTGGGACTA AAGCATAGAC CACCAGGCTG 60

AGTATCCTGA CCTGAGTCAT CCCCAGGGAT CAGGAGCCTC CAGCAGGGAA CCTTCCATTA 120

TATTCTTCAA GCAACTTACA GCTGCACCGA CAGTTGCG ATG AAA GTT CTA ATC TCT 176

Met Lys Val Leu Ile Ser

TCC CTC CTG TTG CTG CCA CTA ATG CTG ATG TCC ATG GTC TCT AGC

Ser Leu Leu Leu Leu Pro Leu Met Leu Met Ser Met Val Ser Ser

-15

-10

-5

AGC CTG AAT CCA GGG GTC GCC AGA GGC CAC AGG GAC CGA GGC CAG GCT

Ser Leu Asn Pro Gly Val Ala Arg Gly His Arg Asp Arg Gly Gln Ala

1 5 10 15

TCT AGG AGA TGG CTC CAG GAA GGC GGC CAA GAA TGT GAG TGC AAA GAT

Ser Arg Arg Trp Leu Gln Glu Gly Gly Gln Glu Cys Glu Cys Lys Asp

20
25
30

TGG TTC CTG AGA GCC CCG AGA AGA AAA TTC ATG ACA GTG TCT GGG
Trp Phe Leu Arg Ala Pro Arg Arg Lys Phe Met Thr Val Ser Gly
35
40
45

(2) INFORMATION FOR SEQ ID NO: 55:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 146 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 99..140
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.2 seq LLLLQLSLPSPTS/SP
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 55:

AAAAATGATG TCACTGGGAA CTGCAGTCAT TTGAAAAGAT AGCAATCAAG CATTTCTTTC 60
AGAGCCCTGT TCATCTTTCA GTGCCTTTGC TTCTCCTG ATG CTT TTG CTC CTT CAA 116

Met Leu Leu Leu Gln
-10

TTA TCT CTG CCT TCT CCC ACC TCC TCT CCG Leu Ser Leu Pro Ser Pro Thr Ser Ser Pro

146

- (2) INFORMATION FOR SEQ ID NO: 56:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 105 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 25..75
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.1

seq LSFKLLLLAVALG/FF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 56:

AGCCCCTGCT GCTCTGGGCA GACG ATG CTG AAG ATG CTC TCC TTT AAG CTG

Met Leu Lys Met Leu Ser Phe Lys Leu

-15

-10

CTG CTG CTG GCC GTG GCT CTG GGC TTC TTT GAA GGA GAT GCT AAG TTT
Leu Leu Leu Ala Val Ala Leu Gly Phe Phe Glu Gly Asp Ala Lys Phe

-5

GGG GAA Gly Glu

105

- (2) INFORMATION FOR SEQ ID NO: 57:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 344 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate

	(:	ix)	(B) (C)	NAME LOCA	E/KE) ATION NTIFI ER IN	1: 1: [CAT]	38: ION !	203 METH(DD: V							
	()	κi) :	SEQUI	ENCE	DESC	CRIP	rion	: SE(QID	NO:	57:					
AGC'	rcct:	rcc ·	TGTT	ccc	rg go	CGGC	CCCT	C GC'	TTCT'	гсст	TCT	GGAT(GGG (GGCC	CAGGG	G 60
GCC	CAGG	AGA (GTATA	AAAG	GC GA	ATGTO	GGAG	G GT	GCCC	GGCA	CAA	CCAG	ACG (CCCA	GTCAC	A 120
GGG	CGGA	GAG (CHST		ATG (Met E	lis A					Met 1					170
ACG Thr	CTT Leu -10	GCC Ala	CTC Leu	CTG Leu	GGG Gly	GRC Xaa -5	MCC Xaa	AMC Xaa	TGG Trp	GCA Ala	GGG Gly 1	AAG Lys	ATG Met	TAT Tyr	GGC Gly 5	218
CCT Pro	GGA Gly	GGA Gly	GG C Gly	AAG Lys 10	TAT Tyr	TTC Phe	AGC Ser	ACC Thr	ACT Thr 15	GAA Glu	GAC Asp	TAC Tyr	GAC Asp	CAT His 20	GAA Glu	266
ATC Ile	ACA Thr	GGG Gly	CTG Leu 25	CGG Arg	GTG Val	TCT Ser	GTA Val	GGT Gly 30	CTT Leu	CTC Leu	CTG Leu	GTG Val	AAA Lys 35	AGT Ser	GTC Val	314
			CTT Leu													344
(2)	INFO	ORMA'	rion	FOR	SEQ	ID 1	NO: :	58:								
	<u>ذ</u>) -	L) SE	(B) (C)	LENC TYPE STRA	CHARA STH: L: NU ANDEE	267 CLEI NESS	base C AC S: DC	e pai CID DUBLE								
	(i	.i) N	40LEC	CULE	TYPE	: CI	ONA									
	7)	7i) (ORGA	SOUF NISM SUE I	1: Hc				state	<u>:</u>					
	(i	ix) i	FEATU (A) (B)	NAME	:/KEY	': si I: 58	.g_pe	eptic	ie							

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 58:

(C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 8

seq VSAVLCVCAAAWC/SQ

ATG CTC AAG GTG TCA GCC GTA CTG TGT GTG TGT GCA GCC GCT TGG TGC Met Leu Lys Val Ser Ala Val Leu Cys Val Cys Ala Ala Ala Trp Cys -15 -5	105
AGT CAG TCT CTC GCA GCT GCC GCG GCG GTG GCT GCA GCC GGG GGG CGG Ser Gln Ser Leu Ala Ala Ala Ala Ala Val Ala Ala Ala Gly Gly Arg 1 5 10 15	153
TCG GAC GGC GGT AAT TTT CTG GAT GAT AAA CAA TGG CTC ACC ACA ATC Ser Asp Gly Gly Asn Phe Leu Asp Asp Lys Gln Trp Leu Thr Thr Ile 20 25 30	201
TCT CAG TAT GAC AAG GAA GTC GGA CAG TGG AAC AAA TTC CGA GAC GAT Ser Gln Tyr Asp Lys Glu Val Gly Gln Trp Asn Lys Phe Arg Asp Asp 35 40 45	249
GAT TAT TTC CGC ACT GGG Asp Tyr Phe Arg Thr Gly 50	267
(2) INFORMATION FOR SEQ ID NO: 59: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 258 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 124174 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 7.8</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 59:	
AAGCATAAGA AGTGATTGAG CCACAAGTAT ACTGAAGGAA GGGCTCCCTC GAGTTGTGGT	60
GTGAAGAGAT AAATCACCAG TCACAGACTA TGCACCCGAC TGCTGCTGTT CAGTCCAGGG	120
AAA ATG AAA GTT GGA GTG CTG TGG CTC ATT TCT TTC ACC TTC ACT Met Lys Val Gly Val Leu Trp Leu Ile Ser Phe Phe Thr Phe Thr -15 -10 -5	168
GAC GGC CAC GGT GGC TTC CTG GGG AAA AAT GAT GGC ATC AAA ACA AAA ASp Gly His Gly Gly Phe Leu Gly Lys Asn Asp Gly He Lys Thr Lys 1 5 10	216
AAA GAA CTC ATT GTG AAT AAG AAA AAA CAT CTA GGC CTC GGG Lys Glu Leu Ile Val Asn Lys Lys His Leu Gly Leu Gly	258

VO 99/06550		PCT/IB98/01232
,	45	

15	20	25

- (2) INFORMATION FOR SEQ ID NO: 60:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 211 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 155..202
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.7

seq ILLDLICLLFITA/CV

211

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 60:

ACTGAAATAG GAAAGTAAGA TTTATACCCA TTATTCAGCC AAAATCTGTT TTTCTTTAAC 60

TTCTACCCAT TGTTCCTAAG TCTGCCCTCT GGGGGCTGTA GAAAATAATG AAGATGATGT 120

TATTAATGAT AACCAGTGCT TGCTGTAACC AGTT ATG TGC ATT ATT TTA TTG GAT 175

Met Cys Ile Ile Leu Leu Asp
-15

TTA ATT TGT TTA CTC TTT ATA ACA GCA TGT GTG GGG
Leu Ile Cys Leu Leu Phe Ile Thr Ala Cys Val Gly
-5

(2) INFORMATION FOR SEQ ID NO: 61:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 316 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 131..307
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.6

seq FMVFGSFFPLISC/QP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 61:

ACATGGATTG ATTTGTTATT TGGGGATTAA ATTAGGCAGG GCACATAGTA GGGCCTCCTT	60
GGATGTTTGA TGGCTGTTGA ATGAACGTAA GTGAATCTGT TCAGTTTTAG GGTTTTATTG	120
CATTTTTGAT ATG GAT TGT GCC AGT ATA TCT GTA AAG TTC ACT TCT ATG Met Asp Cys Ala Ser Ile Ser Val Lys Phe Thr Ser Met -55 -50	169
GCT ACC ATG CAT GAC TTG AGT CAG TTC TGG GCT TCT AGA GGA GAG GTT Ala Thr Met His Asp Leu Ser Gln Phe Trp Ala Ser Arg Gly Glu Val -45 -35	217
ACA AAC TGG TGG CCA GTA GGA CAA ACT AGC CTA CCA CTG TTT TAT TTG Thr Asn Trp Trp Pro Val Gly Gln Thr Ser Leu Pro Leu Phe Tyr Leu -30 -25 -20 -15	265
GCT TTC ATG GTG TTT GGT TCT TTT TTT CCT TTA ATT TCC TGC CAG CCC Ala Phe Met Val Phe Gly Ser Phe Phe Pro Leu Ile Ser Cys Gln Pro -10 -5 1	313
GGG Gly	316

(2) INFORMATION FOR SEQ ID NO: 62:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 317 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 147..206
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.6

seq LVVLFGITAGATG/AK

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 62:

ACTITIGCAC TAGCAGTAGC AAGGAAGGGG GGTGGGCGCT CTTTCTTTTT CTCTTAGAAG 60

AGGGTTTAGC ACAGGTTTTT TCGTTCTCAC TTCCACACCA CCTTACCGCC TCCCGACCCC 120

CCCTCTCCCC CTCCCCACCT ATCGTC ATG ACG GCC TCT CCG GAT TAC TTG GTG Met Thr Ala Ser Pro Asp Tyr Leu Val -20 -15

Val	CTT Leu -10	TTT Phe	GGG Gly	ATC Ile	ACT Thr	GCT Ala ~5	GGG Gly	GCC Ala	ACC Thr	GGG Gly	GCC Ala 1	AAG Lys	CTA Leu	GGC Gly	TCG Ser 5	221
GAT Asp	GAG Glu	AAG Lys	GAG Glu	TTG Leu 10	ATC Ile	CTG Leu	CTG Leu	TTC Phe	TGG Trp 15	AAA Lys	GTC Val	GTG Val	GAT Asp	CTN Leu 20	GCC Ala	269
AAC Asn	AAG Lys	AAG Lys	GTG Val 25	GGA Gly	CAG Gln	TTG Leu	CAC His	GAA Glu 30	BKA Xaa	HGN Xaa	TTA Leu	GAC Asp	CGG Arg 35	ATC Ile	TGG Trp	317
(2)	(2) INFORMATION FOR SEQ ID NO: 63: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 282 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 4690 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 7.6															
					DESC	RIPT	'ION:	SEC	_		LAAA		,			
	(;	(i) S	SEQUE	NCE					10	NO.	63:					
AAGC						GGAC	GCA1	GCC					t Va		GC GTT /s Val	57
CTC	GGCT GTT	rgg 1	CCCC		AG TI	GCA	GGA	GCT	GTG	TTC GCG	TCTC	Me -1 TTC	t Va 5 CTA	1 Cy ATC	/s Val CTG	57 105
CTC Leu CGA	GTT Val -10	CTA Leu	GCT Ala GTA	GGA# GCG	GCC Ala	GCA Ala -5 CGT	GGA Gly TCC	GCT Ala ATG	GTG Val GAC	GCG Ala GTT	GTT Val 1	Me -1 TTC Phe	t Va 5 CTA Leu CGG	ATC Ile	CTG Leu 5	
CTC Leu CGA Arg	GGCT GTT Val -10 ATA Ile	CTA Leu TGG Trp	GCT Ala GTA Val	GCG Ala GTG Val	GCC Ala CTT Leu	GCA Ala -5 CGT Arg	GGA Gly TCC Ser	GCT Ala ATG Met	GTG Val GAC Asp 15	GCG Ala GTT Val	GTT Val 1 ACG Thr	Me -1 TTC Phe CCC Pro	t Va 5 CTA Leu CGG Arg	ATC Ile GAG Glu 20	CTG Leu 5 TCT Ser	105
CTC Leu CGA Arg CTC Leu	GTT Val -10 ATA Ile AGT Ser	CTA Leu TGG Trp ATC Ile	GCT Ala GTA Val TTG Leu 25	GCG Ala GTG Val 10	GCC Ala CTT Leu GTG Val	GCA Ala -5 CGT Arg GCT Ala	GGA Gly TCC Ser GGG Gly	GCT Ala ATG Met TCC Ser 30 AAT	GTG Val GAC Asp 15 GGT Gly	GCG Ala GTT Val GGG Gly TAC	GTT Val 1 ACG Thr CAT His	Me -1 TTC Phe CCC Pro ACC Thr	CTA Leu CGG Arg ACT Thr 35	ATC Ile GAG Glu 20 GAG Glu	CTG Leu 5 TCT Ser ATC Ile	105 153

(2)	INFORMATION	FOR	SEQ	ΙD	NO:	64:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 293 base pairs

60

- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 48..179
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.5

seq LMIPLLLTPITA/TS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 64:

ACAACTCAAG CCAGACAGGC AGCAATTCCA GAGTCGAAAG AGGCCT										CCTT			AAA Lys	56		
ACC Thr	GGG Gly -40	Asp	GGG Gly	GGT Gly	ACT Thr	TTG Leu -35	AGC Ser	ACC Thr	GAG Glu	AGG Arg	ATA Ile -30	GGA Gly	GGG Gly	GCC Ala	GCT Ala	104
CTC Leu -25	CTC Leu	AGC Ser	CTC Leu	CTG Leu	CTG Leu -20	AAG Lys	AGG Arg	ATG Met	AAG Lys	ATG Met ~15	ACT Thr	TTG Leu	ATG Met	ATA Ile	CCC Pro -10	152
TTG Leu	TTG Leu	CTA Leu	TTG Leu	ACA Thr -5	CCT Pro	ATA Ile	ACT Thr	GCG Ala	ACC Thr	TCC Ser	ACT Thr	TCA Ser	AGG Arg 5	TGG Trp	CCC Pro	200
GAG Glu	ATC Ile	GGA Gly 10	GTA Val	GTG Val	GCT Ala	ATC Ile	CGC Arg 15	TCA Ser	CAA Gln	TTG Leu	AGG Arg	GCT Ala 20	TTG Leu	CAT His	ACC Thr	248
TGT Cys	GGT Gly 25	CAG Gln	GAG Glu	CCC Pro	GTG Val	CCA Pro 30	GCT Ala	ATG Met	GGG Gly	TCA Ser	GAA Glu 35	GGG Gly	GCC Ala	GCG Ala		293

(2) INFORMATION FOR SEQ ID NO: 65:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 340 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR

	(ii)	MOLE	CULE	TYP	E: C	DNA									
	(,	vi)	ORIG (A) (F)	INAL ORGA	ANIS	м : н	omo s	Sapie rmal	ens pro:	state	e					
	(:	ix)	(B) (C)	NAMI LOCA	ATION NTIF:	1: 32 [CAT]	210 ION N	OO METHO	D: 1	re 7	Heijr .5 LQLLI					
	(2	ĸi)	SEQUI	ENCE	DES	CRIP	rion	: SE() ID	NO:	65:					
AGT	AGAC	GCT	CGGG	CACC	AG CI	MGCG	GCAA(Leu (TGG A		52
CAG Gln	TTG Leu -15	GGG Gly	CTC Leu	ACT Thr	TTT Phe	CTT Leu -10	CAG Gln	CTC Leu	CTT Leu	CTC Leu	ATC Ile -5	TCG Ser	TCC Ser	TTG Leu	CCA Pro	100
AGA Arg 1	GAG Glu	TAC Tyr	ACA Thr	GTC Val 5	ATT Ile	AAT Asn	GAA Glu	GCC Ala	TGC Cys 10	CCT Pro	GGA Gly	GCA Ala	GAG Glu	TGG Trp 15	AMT Xaa	148
ATC Ile	ATG Met	TGT Cys	CGG Arg 20	GAG Glu	TGC Cys	TGT Cys	GAA Glu	TAT Tyr 25	GAT Asp	CAG Gln	ATT Ile	GAG Glu	TGC Cys 30	GTC Val	TGC Cys	196
CCC Pro	GGA Gly	AAG Lys 35	AGG Arg	GAA Glu	GTC Val	GTG Val	GGT Gly 40	TAT Tyr	ACC Thr	ATC Ile	CCT Pro	TGC Cys 45	TGC Cys	AGG Arg	AAT Asn	244
GAG Glu	GMG Xaa 50	AAT Asn	GAG Glu	TGT Cys	GAC Asp	TCC Ser 55	TGC Cys	CTG Leu	ATC Ile	CAC His	CCA Pro 60	GGT Gly	TGT Cys	ACC Thr	ATC Ile	292
TTT Phe 65	GAA Glu	AAC Asn	TGC Cys	AMG Xaa	AGC Ser 70	TGC Cys	CGM Arg	AAT Asn	GGC Gly	TCA Ser 75	TGG Trp	GGG Gly	GGT Gly	ACC Thr	TTG Leu 80	340
(2)			rion													
	(<u>i</u>	.) SI	(B)	ICE C LENG TYPE STRA	TH: : NU	351 CLEI	base C AC	pai								

- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE: -

 - (A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:

(A)	NAME/KEY:	sig_peptide
		ara-beherme

- (B) LOCATION: 112..192
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 7.2 seq SLLFFLLLEGGXT/EQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 66:

															VACGGA	60
GAA	GGCV	AKK	RCNN	NNRC	TT G	AAGG	TTCT	G TC	ACCT	TTTG	CAG	TGGT	CCA		G AGA t Arg	117
RAA Xaa -25	гÀг	TGG Trp	AAA Lys	ATG Met	GGA Gly -20	GGC Gly	ATG Met	AAA Lys	TAC Tyr	ATC Ile -15	TTT Phe	TCG Ser	TTG Leu	TTG Leu	TTC Phe -10	165
TTT Phe	CTT Leu	TTG Leu	CTA Leu	GAA Glu -5	GGA Gly	GGC Gly	KAA Xaa	ACA Thr	GAG Glu 1	CAA Gln	GTR Val	AMN Xaa	CAT His 5	TCA Ser	GAG Glu	213
ACA Thr	TAT Tyr	TGC Cys 10	ATG Met	TTT Phe	CAA Gln	GAC Asp	AAG Lys 15	AAG Lys	TAC Tyr	AGA Arg	GTG Val	GGT Gly 20	GAG Glu	AGA Arg	TGG Trp	261
CAT His	CCT Pro 25	TAC Tyr	CTG Leu	GAA Glu	CCT Pro	TAT Tyr 30	GGG Gly	TTG Leu	GTT Val	TAC Tyr	TGC Cys 35	GTG Val	AAC Asn	TGC Cys	ATC Ile	309
TGC Cys 40	TCA Ser	GAG Glu	RAT Xaa	GGG Gly	AAT Asn 45	GTG Val	CTT Leu	TGC Cys	AGC Ser	CGA Arg 50	GTC Val	AGA Arg	TGT Cys			351

(2) INFORMATION FOR SEQ ID NO: 67:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 310 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 68.124
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.2

seq VSIMLLLVTVSDC/AV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 67:

							•			
AGTGA			AGA Arg							109
ACT G Thr V -5										157
CAG T Gln C										205
CTG C										253
GGC A										301
CCT T	GC	TTG								310

(2) INFORMATION FOR SEQ ID NO: 68:

Pro Cys Leu

60

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 380 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 240..302
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.2

seq SALLFSLLCEAST/VV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 68:

ACCTTTCTGG ACGTTGCAAA CTGTGACATA TAAAAGCTGT TAGCTGCTCC	TCTAGCCAGC	60
AGCATTCAAA CCTTGCAGAG CTTTGCTCTC AGAGAGTTTG TAAAAAGACA	CACTCCTCTT	120
ACAAGAGTTC ATGCTACCAC ATAGCAAAGA ACCTTAAATT TTTGGAAGAA	CAATATATTC	180
ATTTTGGCAT TGTGCAGAGC AAAGTAAACT CGGTGGCCTC TTCTTCTCCA	CCCCTCAAR	239
ATG ATA GCR ATC TCT GCC GTC AGC AGT GCA CTC CTG TTC TC Met Ile Ala Ile Ser Ala Val Ser Ser Ala Leu Leu Phe Se		287

WO 99/06550 PCT/IB98/01232

	-20					-15					-10					
TGT Cys -5	GAA Glu	GCA Ala	AGT Ser	ACC Thr	GTC Val 1	GTC Val	CTA Leu	CTC Leu	AAT Asn 5	TCC Ser	ACT Thr	GAC Asp	TCA Ser	TCC Ser 10	CCG Pro	335
CSA Xaa	ACC Thr	AAT Asn	AAT Asn 15	TTC Phe	RCT Xaa	GAT Asp	AWT Xaa	GAA Glu 20	GCA Ala	GCT Ala	CTG Leu	AAA Lys	GCA Ala 25	CAT His		380

(2) INFORMATION FOR SEQ ID NO: 69:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 435 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 181..243
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.2

seq SALLFSLLCEAST/VV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 69:

AGC	ATTC.	AAA	CCTT	GCAG.	AG C	TTTG	CTCT	C AG	AGAG	TTTG	TAA	AAAG	ACA	CACT	ССТСТТ	60
ACA	AGAG	TTC .	ATGC	TACC	AC A	TAGC	AAAG.	A AC	CTTA	AATT	TTT	GGAA	GAA	CAAT.	ATATTC	120
MAT	TTTG	GCA	TTGT	GCAG	AG C	AAAG'	TAAA	C TC	GGTG	GCCT	CTT	CTTC	TCC .	ACCC	CTCAAA	180
ATG Met	ATA Ile -20	GCA Ala	ATC Ile	TCT Ser	GCC Ala	GTC Val -15	AGC Ser	AGT Ser	GCA Ala	CTC Leu	CTG Leu -10	TTC Phe	TCC Ser	CTT Leu	CTC Leu	228
TGT Cys -5	GAA Glu	GCA Ala	AGT Ser	ACC Thr	GTC Val 1	GTC Val	CTA Leu	CTC Leu	AAT Asn 5	TCC Ser	ACT Thr	GAC Asp	TCA Ser	TCC Ser 10	CCG Pro	276
CCA Pro	ACC Thr	AAT Asn	AAT Asn 15	TTC Phe	ACT Thr	GAT Asp	ATT Ile	GAA Glu 20	GCA Ala	GCT Ala	CTG Leu	AAA Lys	GCA Ala 25	CAA Gln	TTA Leu	324
GAT Asp	TCA Ser	GCG Ala 30	GAT Asp	ATC Ile	CCC Pro	AAA Lys	GCC Ala 35	AGG Arg	CGG Arg	AAG Lys	CGC Arg	TAC Tyr 40	ATT Ile	TCG Ser	CAG Gln	372
AAT Asn	GAC Asp 45	ATG Met	ATC Ile	GCC Ala	ATT Ile	CTT Leu 50	GAT Asp	TAT Tyr	CAT His	AAT Asn	CAA Gln	GTT Val	CGG Arg	GGC Gly	AAA Lys	420

GTG	TTC	CCA	MCG	GCA
Val	Phe	Pro	Xaa	Ala
60				

(2) INFORMATION FOR SEQ ID NO: 70:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 426 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide(B) LOCATION: 352..417

 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.2 seq LLTLVLCVAVAYE/RO
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 70:

ATTGAGCTGT CTGCAGCAGA GCTGAGAGGA CCAGCCATTT TACTTATGGA AAACAGTGTG	60
GCATATTCTG CTGAGCTTCG CCCTGGAAGA AGCCTCTTTT ATACATCTCT TCAGGGAAGA	120
GAGAAGCAAT GGGCATGTTA GTATACAATG ATCACAGCCA CGCAGGCCTG CAAGCTGCCT	180
TTTGGACAGG CTGTTGACTG CCGTTCCAAT TAGCTGATTG GAGAATGTGG AATGCAGAGT	240
GATAATGCTG CATATCTGCT ATCAGGCAGC AGCAAAGGTT TTTGTCTTGG GAAGGCAAGC	300
TTTCCCTGCA ATATTATCTC AGCAGCTCCC TAGCTGCTTA CCCTGAAAAC G ATG GAT Met Asp	357
CCA AAC GGA GGG TGT TGC ACT CTG CTA ACG CTG GTC CTG TGC GTG GCT Pro Asn Gly Gly Cys Cys Thr Leu Leu Thr Leu Val Leu Cys Val Ala -20 -15 -10 -5	405
GTG GCA TAT GAG CGG CAG GAG Val Ala Tyr Glu Arg Gln Glu	426

(2) INFORMATION FOR SEQ ID NO: 71:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 389 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR

WO 99/0655		54	PCT/IB98/01232
(ii) M	OLECULE TYPE: CDNA		
	RIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Cancerous	prostate	
		Von Heijne matrix ore 7.2 LFTFSTSLPSSLS/SS	
(xi) SI	EQUENCE DESCRIPTION: SEQ ID) NO: 71:	
ACAATACCTG TT	FACTTATAT ACTTTTCTTT GTCTAA	AAAA GAAATAAGAT CTGTCTA	GAT 60
	ITAGGGAGA TTCTGATTAA CAGAAT		
	ГАТАТТТТБ ТАССААТТТА ТАТААА		
	GCTAGAGAT GAAATATCGT TTTCCT		
CACCCCATGA AC	CTCATACAG GTCAGAATTA AACCCC	GAGC TTTGTTT ATG GAG GGT Met Glu Gly -25	
-20	TTC CAA GTA TTT CTT TCT CTT Phe Gln Val Phe Leu Ser Leu -15	Phe Thr Phe Ser Thr Ser	A 344
TTA CCA TCA T Leu Pro Ser S -5	CA TTG TCG TCA TCA TTG er Leu Ser Ser Ser Leu l	TCA TCA TCC AAT GGG Ser Ser Ser Asn Gly 5	389
(2) INFORMATI	ON FOR SEQ ID NO: 72:		
() (1	UENCE CHARACTERISTICS: A) LENGTH: 328 base pairs B) TYPE: NUCLEIC ACID C) STRANDEDNESS: DOUBLE		

(2

- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 194..316
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7

seq FLCMLAAIDLALS/TS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 72:

·.

ATGAGTCAGC CTGAAAGGAA CAGGCCGAAC TGCTGTATGG GCTCTACTGC CAGTGTGACC	60
TCACCCTCTC CAGTCACCCC TCCTCAGTTC CAGCTATGAG TTCCTGCAAC TTCACACATG	120
CCACCTTTGT GCTTAATKGG AATCCCAGGG ATTAGAGAAA GCCCATTTCT GGGTTGGCTT	180
CCCCCTCCTT TCC ATG TAT GTA GTG GCA ATG TTT GGA AAC TGC ATC GTG Met Tyr Val Val Ala Met Phe Gly Asn Cys Ile Val -40 -35 -30	229
GTC TTC ATC GTA AGG ACG GAA CGC AGC CTG CAC GCT CCG ATG TAC CTC Val Phe Ile Val Arg Thr Glu Arg Ser Leu His Ala Pro Met Tyr Leu -25 -15	277
TTT CTC TGC ATG CTT GCA GCC ATT GAC CTG GCC TTA TCC ACA TCC ACC Phe Leu Cys Met Leu Ala Ala Ile Asp Leu Ala Leu Ser Thr Ser Thr -10 -5 1	325
ATG Met	328
(2) INFORMATION FOR SEQ ID NO: 73:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 267 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR 	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide - (B) LOCATION: 79207 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 7 seq PWFLAPWCPGTQS/NR</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 73:	
ACCCTTCGTT CTGGTTCTGG TTCTAGTTCT GGTTCTAACA ACTCACAATC CCTTTAGCTT	60
TCTCTCCCCT CCCTTTGA ATG AGA GAA ACT AMC CCG CTT CCG AAG CCC CTG Met Arg Glu Thr Xaa Pro Leu Pro Lys Pro Leu -40 -35	111
AAA GAC ACT GCT CCT TCC TCT CAT GGA GTT GGC TCC GAC AGC CCG TCT Lys Asp Thr Ala Pro Ser Ser His Gly Val Gly Ser Asp Ser Pro Ser -30 -25 -20	159
GCC ACC AGG CCA TGG TTC CTT GCC CCA TGG TGT CCT GGG ACC CAG AGC Ala Thr Arg Pro Trp Phe Leu Ala Pro Trp Cys Pro Gly Thr Gln Ser	207

WO 99/0	6550			PCT/IB98/0123:	2
-15		-10	56 -5		
AAC AGG AT	ר דהד ראר רכא		_		
Asn Arg II	e Cys His Pro	Pro Leu Ser	TCT CCC CCA GAT Ser Pro Pro Asp 10	CAA GCG ACG 255 Gln Ala Thr 15	
TGC CTC AG Cys Leu Ar				267	
(2) INFORM	ATION FOR SEQ	ID NO: 74:			
(i)	(B) TYPE: NO	301 base pai CLEIC ACID NESS: DOUBLE			
(ii)	MOLECULE TYPE	: CDNA			
(vi)	ORIGINAL SOUP (A) ORGANISM (F) TISSUE T	: Homo Sapie	ns prostate		
(ix)	FEATURE: (A) NAME/KEY (B) LOCATION (C) IDENTIFI (D) OTHER IN	: 23202 CATION METHO FORMATION:	e D: Von Heijne ma score 7 seq VLVVLALRSLGR		
(xi)	SEQUENCE DESC	RIPTION: SEQ	ID NO: 74:		
AAGTGAGGCT	TGGAAAGGCG TC	ATG GAC AGA Met Asp Arg	CCT GGG TCG CTT Pro Gly Ser Leu	TCT GTC TTC 52 Ser Val Phe	

GGG TCC CTC CCG GCT TCG CTC GGG ACC TGG CTC TCA AGC CCA GCT TGG

Gly Ser Leu Pro Ala Ser Leu Gly Thr Trp Leu Ser Ser Pro Ala Trp

CTG GTG GAC AGA CCG GTG CGC TCT GCA CAC CCG AGT GCG AAT TCC ACC Leu Val Asp Arg Pro Val Arg Ser Ala His Pro Ser Ala Asn Ser Thr

GGC GTG AGA ATG AGC GTG CTC GTG GTC CTG GCC CTG AGG TCC CTG GGT Gly Val Arg Met Ser Val Leu Val Val Leu Ala Leu Arg Ser Leu Gly

CGC AGC TGT TCC CTC TCC CAG GCT GCC CCC TCC AGG TGG ACG CGG TCA

Arg Ser Cys Ser Leu Ser Gln Ala Ala Pro Ser Arg Trp Thr Arg Ser

AAC GAT GCC CCG CAG CCT CCT GGG TCT CAG CAC ATA TTC CAC ACC TAH Asn Asp Ala Pro Gln Pro Pro Gly Ser Gln His Ile Phe His Thr Xaa

-30

-15

1

GTS CCC GGG

-40

25

-50

100

148

196

244

301

Val Pro Gly

(ix) FEATURE:

(A) NAME/KEY: sig_peptide (B) LOCATION: 160..216

(D) OTHER INFORMATION: score 7

(C) IDENTIFICATION METHOD: Von Heijne matrix

(2) INFORMATION FOR SEQ ID NO: 75:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 110 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 365 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 7</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 75:	
AT ATG CAT TAT TTT GTT GCT GGG AAA GTA ATC CTT CTC TCT TAT Met His Tyr Phe Val Ala Gly Lys Val Ile Leu Leu Phe Ser Tyr -20 -15 -10	47
CCA TCA TGT TGT TTG TGT TTC TTG GTG TAC AGG AGA GTA AGC WAT TTA Pro Ser Cys Cys Leu Cys Phe Leu Val Tyr Arg Arg Val Ser Xaa Leu -5 1 5 10	95
TTT AAG TGC TTT GAG Phe Lys Cys Phe Glu 15	110
(2) INFORMATION FOR SEQ ID NO: 76:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 318 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: DOUBLE(D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Cancerous prostate	

seq STVVLQVLTQATS/QD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 76:

AGACGCCARA CATGGCGTGT TCCTAGAAGC CGCTTTCGGC ATCAGTAGGC GGCGGCGTGG GGTCTGGCAK CGTGGGGAGA GGGAMCAACC GACGCCACTT CGTGTTGGGA AGTGGGAGCG GGANRGCCGG GCAATTCCCG ACCGAACCAA ACGGTTTCC ATG GAT CTC AAT AGT 174 Met Asp Leu Asn Ser GCC AGC ACT GTT GTT CAG GTG TTA ACA CAG GCC ACC AGT CAG GAT 222 Ala Ser Thr Val Val Leu Gln Val Leu Thr Gln Ala Thr Ser Gln Asp -10 ACT GCT GTG TTA AAA CCA GCT GAG GAG CAG TTG AAG CAG TGG GAG ACA Thr Ala Val Leu Lys Pro Ala Glu Glu Gln Leu Lys Gln Trp Glu Thr 270 CAG CCA GGW TTC TAT TCA GTG TTG CTG AAT ATT TTC ACC AAC CAC GGG 318 Gln Pro Gly Phe Tyr Ser Val Leu Leu Asn Ile Phe Thr Asn His Gly 25

(2) INFORMATION FOR SEQ ID NO: 77:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 325 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 95..313
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7

seq FLCMLAAIDLALS/TS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 77:

ATGAGTCAGC CTGAAAGAAC AGGCCGAACT GCTGTATGGG CTCTACTGCC AGTGTGACCT 60

CACCCTCTCC AGTCACCCCT CCTCAGTTCC AGCT ATG AGT TCC TGC AAC TTC ACA 115

Met Ser Ser Cys Asn Phe Thr

CAT GCC ACC TTT GTG CTT ATT GGT ATC CCA GGA TTA GAG AAA GCC CAT
His Ala Thr Phe Val Leu Ile Gly Ile Pro Gly Leu Glu Lys Ala His
-65 -55

									5	9						
TTC Phe -50	TGG Trp	GTT Val	GGC Gly	TTC Phe	CCC Pro -45	CTC Leu	CTT Leu	TCC Ser	ATG Met	TAT Tyr -40	GTA Val	GTG Val	GCA Ala	ATG Met	TTT Phe -35	211
GGA Gly	AAC Asn	TGC Cys	ATC Ile	GTG Val -30	GTC Val	TTC Phe	ATC Ile	GTA Val	AGG Arg -25	ACG Thr	GAA Glu	CGC Arg	AGC Ser	CTG Leu -20	CAC His	259
GCT Ala	CCG Pro	ATG Met	TAC Tyr -15	CTC Leu	T TT Phe	CTC Leu	TGC Cys	ATG Met -10	CTT Leu	GCA Ala	GCC Ala	ATT Ile	GAC Asp -5	CTG Leu	GCC Ala	307
				ACC Thr												325
(2)	INFO	RMAT	CION	FOR	SEQ	ID t	10: 7	78:								
	(i	.) SE	(A) (B) (C)	ICE C LENG TYPE STRA TOPC	TH: : NU NDED	415 CLEI NESS	base C AC C DC	e pai CID OUBLE								
	(i	i) M	OLEC	ULE	TYPE	: CI	NA									
	(v	ri) C	(A)	NAL ORGA TISS	NISM	: Ho				rost	ate					
	(i	×) F	(A) (B) (C)	JRE: NAME LOCA IDEN OTHE	TION TIFI	: 17 CATI	93 ON M	846 IETHO IN:	D: V	on H e 6. PLFF	9					
	(×	i) S	EQUE	NCE	DESC	RIPT	'ION:	SEÇ) ID	NO:	78:					
ACAA	AATC	AA G	AAA	ATCC	A CA	TAGA	TGGT	CAA	AAT <i>F</i>	ATTC	ATAG	GTGA	CT C	SAGAG	STATCC	60
aaa1	GGGC	CA G	GTGA	ACTGA	AG AA	TAC	CAA	A CAG	GCCA	AGAA	TAAT	'ATCI	GT C	STTAA	LATTTG	120
ACCC	TCTA	TT T	TATI	AAC	T AT	CTG	CATO	ACC	TTTC	CTCT	GTAC	CTGC	TG I	agt <i>i</i>	ACTC	178
ATG Met	TAT Tyr -55	AGA Arg	CTC Leu	AGT Ser	CTT Leu	ATA Ile -50	GCA Ala	GGC Gly	CCT Pro	GGG Gly	TCC Ser -45	TAT Tyr	CCT Pro	GTG Val	CTA Leu	226
AGA Arg -40	TGG Trp	GGA Gly	GTT Val	TGG Trp	GAC Asp -35	ATC Ile	CCT Pro	AGT Ser	TCA Ser	TTA Leu -30	GTT Val	CAA Gln	GTG Val	ACT Thr	TAC Tyr -25	274

CAT CAG CCC AAC CTC ACT ACA AAT TTG GAT CTG CCT TTG TTC TTC AGT

His Gln Pro Asn Leu Thr Thr Asn Leu Asp Leu Pro Leu Phe Phe Ser

TGT AGT ATC TCG GCT ACC CAT TCT TGT GTC AAG CCT CCA TCT GTA ATT

-15

-20

322

370

WO 99/065	50		60		PC	T/IB98/012
Cys Ser Ile	Ser Ala Th	r His Ser	Cys Val	Lys Pro Pro	Ser Val Ile	
ATT GGT ATC Ile Gly Ile 10	TCT TCT TT Ser Ser Ph	C CTG AGC e Leu Ser 15	TTT CCT Phe Pro	TAT CAA ACT Tyr Gln Thr 20	TTG GTA Leu Val	415
(2) INFORMA	TION FOR SEC	Q ID NO: 7	9:			
(i) SI	EQUENCE CHAP (A) LENGTH: (B) TYPE: N (C) STRANDE (D) TOPOLOG	400 base NUCLEIC AC DNESS: DO	pairs ID			
(ii) N	MOLECULE TYP	E: CDNA				
(vi) (ORIGINAL SOU (A) ORGANIS (F) TISSUE	M: Homo Sa	apiens mal prost	ate		
	CEATURE: (A) NAME/KE (B) LOCATIO (C) IDENTIF (D) OTHER I	N: 12819 ICATION ME NFORMATION	99 ETHOD: Vo N: score seq L	6.9 CFLLLAVAMSE	•	
(x1) S	EQUENCE DES	CRIPTION:	SEQ ID N	10: 79:		
					CA AACAGCCCTC	
					TC CTAGTCCAGG	120
CATCCCG ATG Met	TTG GTG GAT Leu Val Asp	GGC CCA T Gly Pro S -20	CT GAG C Ser Glu A	GG CCA GCC arg Pro Ala -15	CTG TGC TTC Leu Cys Phe	169
TTG CTG TTG Leu Leu Leu -10	GCT GTG GCA Ala Val Ala -5	ATG TCT T Met Ser P	TTC TTC G Phe Phe G	GC TCA GCT ly Ser Ala	CTA TCC ATA Leu Ser Ile 5	217
GAT GAA ACA Asp Glu Thr	CGG GCG CAT Arg Ala His 10	CTG TTG T Leu Leu L	TG AAA G eu Lys G 15	AA AAG ATG lu Lys Met	ATG CGG CTG Met Arg Leu 20	265
GGG GGG CGG Gly Gly Arg 25	CTG GTG CTG Leu Val Leu	AAC ACC A Asn Thr L 30	AG GAG G ys Glu G	AG CTG GCC lu Leu Ala 35	AAT GAG AGG Asn Glu Arg	313
CTC ATG ACG Leu Met Thr 40	CTC AAA ATC Leu Lys Ile	GCT GAG A Ala Glu M 45	TG AAG G let Lys G	AG GCC ATG lu Ala Met 50	AGG ACC CTG Arg Thr Leu	361
		•		•		

ATA TTC CCA CCC AGC ATG CAC TTT TTC CAG GCC AAA TGG Ile Phe Pro Pro Ser Met His Phe Phe Gln Ala Lys Trp

65

60

400

WO 99/06550 61

(2)	INF	ORMA	TION	FOR	SEQ	ID i	NO: 1	30:						
	i)	i) Si	(B) (C)	NCE (LENC TYPE STRA	TH: : NU ANDEC	212 ICLEI NESS	base IC AC S: DC	e pai CID DUBLE						
	i)	Li) I	MOLE	CULE	TYPE	E: CI	ANC							
	7)	7i) (NAL ORGA TISS	NISM	1: Hc		_		rost	ate			
	(i	(x)	(B) (C)	JRE: NAME LOCA IDEN OTHE	TION TIFI	1: 33 CATI	313 ION N	37 1ETHO	D: V	e 6.	9	atri> ⟨G/H¢		
	()	(i)	SEQUE	ENCE	DESC	RIPT	rion:	SE() ID	NO:	80:			
AACO	CGGC	CCG	CGCC	CCGC	CA TO	GGAG	GACC:	r GG				CTG Leu		53
			CCT Pro -25											101
			CTK Leu											149
			GGG Gly											197
			GTA Val											212
(2)	INFO	ORMA'	TION	FOR	SEQ	ID I	NO: (31:						
	()	i) Si	(B) (C)	LENC TYPE STRA	STH: E: NU ANDEE	269 ICLEI INESS	base C AC S: DC	e pai CID OUBLE						
	()	Li) i	MOLE	CULE	TYPE	E: CI	ANC							
	(7	/i) (INAL ORGA TISS	NISM	i: Ho				state	e			

			(B) (C) (D)	NAM LOC IDE OTH	ATIO NTIF ER I	N: 1 ICAT NFOR	51 ION MATI	METHON:	OD: sco seq	re 6 LFL	.8 FLTS					
	(:	XI)	SEQU	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	81:					
ACC	CTGT	кст	тктс	ATG Met	GTT Val -40	DTC Xaa	TGG Trp	CTC Leu	GTC Val	TTA Leu -35	TTT Phe	GCT Ala	CTT Leu	CAG Gln	ATT Ile -30	50
TAC Tyr	TCC Ser	TAT Tyr	KKY Xaa	AGT Ser -25	ACT Thr	CGA Arg	GAT Asp	CAG Gln	CCT Pro -20	GCA Ala	TCA Ser	CGT Arg	GAK Xaa	AGG Arg -15	CTT Leu	98
CTT Leu	TTC Phe	CTT Leu	TTT Phe -10	CTG Leu	ACA Thr	AGT Ser	ATT Ile	GCG Ala -5	GAA Glu	TRC Xaa	T GC Cys	AGC Ser	ACT Thr 1	CCT Pro	TAC Tyr	146
TCT Ser	CTT Leu 5	TTG Leu	GGT Gly	TTK Xaa	GTC Val	TTC Phe 10	ACG Thr	GTT Val	TCT Ser	TTT Phe	GTT Val 15	GCC Ala	TTG Leu	GGT Gly	GTT Val	194
CTC Leu 20	ACA Thr	CTC Leu	TGC Cys	AAG Lys	TTT Phe 25	TAC Tyr	TTG Leu	CAG Gln	GGT Gly	TAT Tyr 30	CGA Arg	GCT Ala	TTC Phe	ATG Met	AAT Asn 35	242
GAT Asp	CCT Pro	GCC Ala	ATG Met	AAT Asn 40	CGG Arg	GGA Gly	GGT Gly	GCG Ala								269
(2)			(B) (C)	ICE C LENG TYPE STRA	HARA TH: NU	CTER 68 b CLEI NESS	ISTI ase C AC : DO	CS: pair ID UBLE								
	(i	i) M	OLEC				NEAR NA	•								
			RIGI													

- (F) TISSUE TYPE: Hypertrophic prostate
 (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide

(A) ORGANISM: Homo Sapiens

- (B) LOCATION: 9..62
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 6.7 seq LPLLXXXSLPVGA/WL
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 82:

(2) INFORMATION FOR SEQ ID NO: 83:

(i) SE(QUENCE	CHARACTERISTICS:
---------	--------	------------------

- (A) LENGTH: 407 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 258..368
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.7

seq ILYILWYCSVCSS/GS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 83:

AAGGTTGGTC TGGACCGGAA GCGAAGATGG CGACTTCTGG CGCGGCCTCG GCGGASTGGT GATCGGCTGG TGCATATTCG GCCTCTTACT ACTGGCKATT TTGGCATTCT GCTGGATATA TGTTCGTAAA TACCAAAGTC GGCGGGAAAG TGAAGTTGTC TCCACCATAA CAGCAATTTT 180 TTCTCTAGCA ATTGCACTTA TCACATCAGC ACTTCTACCA GTGGATATAT TTTTGGTTTC 240 TTACATGAAA AATCAAA ATG GTA CAT TTA AGG ACT GGG CTA ATG CTA ATG 290 Met Val His Leu Arg Thr Gly Leu Met Leu Met -35 TCA GCA GAC AGA TTG AGG ACA CTG TAT TAT ACG GTT ACT ATA CTT TAT 338 Ser Ala Asp Arg Leu Arg Thr Leu Tyr Tyr Thr Val Thr Ile Leu Tyr ATT CTG TGG TAT TGT TCT GTG TGT TCT TCT GGA TCC CTT TTG TCT ACT 386 Ile Leu Trp Tyr Cys Ser Val Cys Ser Ser Gly Ser Leu Leu Ser Thr -10 -5 TCT ATT ATG AAG AAA AGG ATG 407 Ser Ile Met Lys Lys Arg Met 10

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 348 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Cancerous prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 196240 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.7</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 84:	
AAAAAATTGG TCCCAGTTTT CACCCTGCCG CAGGGCTGGC TGGGGAGGGC AGCGGTTTAG	60
ATTAGCCGTG GCCTAGGCCG TTTAACGGGG TGACACGAGC HTGCAGGGCC GAGTCCAAGG	120
CCCGGAGATA GGACCAACCG TCAGGAATGC GAGGAATGTT TTTCTTCGCA GTGTATAGA	180
GCACACAGAC AGACC ATG GGG ATT CTG TCT ACA GTG AGA	231
GCC AGA GCC CTG GAC GGC TGC AGA AAT GGC ATT GCC CAC CCT GCA AGT Ala Arg Ala Leu Asp Gly Cys Arg Asn Gly Ile Ala His Pro Ala Ser 1 5 10	279
GAG AAG CAC AGA CTC GAG AAA TGT AGG GAA CTC GAG AGC AGC CAC TCG Glu Lys His Arg Leu Glu Lys Cys Arg Glu Leu Glu Ser Ser His Ser 15 20 25	327
GCC CCA-GGA TCA ACC CAG CAG Ala Pro Gly Ser Thr Gln Gln 30 35	348
(2) INFORMATION FOR SEQ ID NO: 85:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 146 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate</pre>	

(ix) FEATURE:

 (A) NAME/KEY: sig_peptide (B) LOCATION: 45113 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.5 seq LTFLQXLLISSLX/RE 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 85:	
ACTCTCCCTC CCCAGTAGAC GCTCGGGCAC CAGCCGCGGC AAGG ATG GAG CTG GGT Met Glu Leu Gly -20	56
TGC TGG ACG CAG TTG GGG CTC ACT TTT CTT CAG STC CTT CTC ATC TCG Cys Trp Thr Gln Leu Gly Leu Thr Phe Leu Gln Xaa Leu Leu Ile Ser -15 -10 -5	104
TCC TTG CHA AGA GAG TAC ACA GTC ATT AAT GAA GCH CGC AAG Ser Leu Xaa Arg Glu Tyr Thr Val Ile Asn Glu Ala Arg Lys 1 5 10	146
(2) INFORMATION FOR SEQ ID NO: 86: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 308 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 201266 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.4 seq FLLCXSVFTDCKG/DV</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 86:	
ACAGAATCAC GTTTTDAGTT GTGCGTGTGC GCGCACACGM GTGTAAAMAG CACTTTCGAT	60
TGTGCCTCCT GTTTTCTCGA GTGGGGACAC TTTAACTACA GTTTASACCT CGGGCGCATM	120
AAGTTTKTCT TCTCTTTCTC TCTGGTTRTT TCTGTTTCTG AGTGGACCAA CAGCAGARCC	180
CACGAGGAKT TGTTTTGAGT ATG GAG CTG TTG CGG GTD TGC TCC TTT TTC TTG Met Glu Leu Leu Arg Val Cys Ser Phe Phe Leu -20 -15	233
CTT TGC TSC TCA GTT TTT ACA GAC TGT AAA GGA GAT GTG TTG TGT GTG Leu Cys Xaa Ser Val Phe Thr Asp Cys Lys Gly Asp Val Leu Cys Val +10 -5 1 5	281

				GCT Ala	

- (2) INFORMATION FOR SEQ ID NO: 87:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 289 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 203..268
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.3 seq TWFLLLPPGQCRA/VG
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 87:

AGAATCTCAC GAGAGAAGAA AACCAGCCAC ATAAAGGATT TGAAAGCTCA ACTTGCTTTC	60
CCACTCTGTT ATCCCTGGAG TTGGCTTGGA TTCACCCTGA AGCCTTCCCC CTCCCGGGGA	120
AAGTTGCTTC ACGTTGCAGC TCAGCAGGTT TGTCCAGCTA CATAGGCTCC AGAAAACAAG	180
AAGCAAGACT GGAAAGCTGG GG ATG ATT GTA CGC CCT CGC CTG AAT CTT ACG Met Ile Val Arg Pro Arg Leu Asn Leu Thr -20 -15	232
TGG TTC CTC CTT CTT CCA CCT GGC CAG TGC AGA GCC GTG GGT GCC ACG Trp Phe Leu Leu Pro Pro Gly Gln Cys Arg Ala Val Gly Ala Thr -10 -5	280
TGG CCC GGG Trp Pro Gly	289

- (2) INFORMATION FOR SEQ ID NO: 88:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 120 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA

(vi)	ORIGINAL	SOURCE:	

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 1..57
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 6.3

seq MVALCCCLWKISG/CE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 88:

ATG CAA TTC TTG TTT AAG ATG GTG GCC TTA TGC TGT TGT CTC TGG AAG

Met Gln Phe Leu Phe Lys Met Val Ala Leu Cys Cys Cys Leu Trp Lys

-15

-10

48

ATC TCC GGC TGT GAG GAA GTC CCT CTA ACT TAC AAC CTG CTC AAG TGC

1 Ser Gly Cys Glu Glu Val Pro Leu Thr Tyr Asn Leu Leu Lys Cys

1 5 10

CTC CTA GAT AAA GCG CAC GTA GGG
Leu Leu Asp Lys Ala His Val Gly
15 20

(2) INFORMATION FOR SEQ ID NO: 89:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 247 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 50..112
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.3

seq CVCAAAXXSQSLX/XX

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 89:

AAAGCGTCCT ATCCGGAGCC AACTGTAGCT GGGATCCAGC GAGAGGAAG ATG CTC AAG 58

Met Leu Lys

GTG TCA GCC GTA CTG TGT GTG TGT GCA GCC GCT TDG TGS AGT CAG TCT

Val Ser Ala Val Leu Cys Val Cys Ala Ala Ala Xaa Xaa Ser Gln Ser

-15

-10

-5

CTC GSM RCT KCC GCG GCG GTG GCT GCA GCC GGG GGG CGG TCG GAC GGC 154

68

Leu Xaa Xaa Xaa Ala Ala Ala Val Ala Ala Ala Gly Gly Arg Ser Asp Gly

GGT AAT TTT CTG GAT ASP ASP Lys Gln Trp Leu Thr Xaa Ile Ser Gln Tyr 30

GAC AAG GAA KTC GGM MAG TGG AAC AAA TTC CGA GAC GAT KAT TAT ASP Lys Glu Xaa Gly Xaa Trp Asn Lys Phe Arg Asp Asp Xaa Tyr 45

(2) INFORMATION FOR SEQ ID NO: 90:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 294 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 124..186
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.3

seq MVALCCCLWKISG/CE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 90:

AAGACGCTGC CTTTAGGGAG AGATAAAAAG CATAATGACA TTAGCTAGGA AAGTTAATTT 60 TCAGTTCTTA CTGAAGTGCT GTATGAAACT GAAATTTCCA AGGAACTGAA TTTTGTGAGC CAR ATG AGC ATG CAR TTC TTG TTT AAG ATG GTG GCC TTA TGC TGT TGT Met Ser Met Gln Phe Leu Phe Lys Met Val Ala Leu Cys Cys Cys CTC TGG AAG ATC TCC GGC TGT GAG GAA GTC CCT CTA ACT TAC AAC CTG Leu Trp Lys Ile Ser Gly Cys Glu Glu Val Pro Leu Thr Tyr Asn Leu 216 CTC AAG TGC CTC CTA GAT AAA GCG CAC TGT GTA CTC CTG ACA CCT TGT Leu Lys Cys Leu Leu Asp Lys Ala His Cys Val Leu Leu Thr Pro Cys 15 GGT TAC ATC TTT TCC TTG ATC AGT CCA GGG 294 Gly Tyr Ile Phe Ser Leu Ile Ser Pro Gly 30 35

(2) INFORMATION FOR SEQ ID NO: 91:

 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 173 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR 	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 114164 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.2</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 91:	
AATTCTTATA GGTGTGTCCA GCAGGCAGTG GCTTGTAGCT GTTCCTTCAG CCACTTAACA	60
GGTTTGATTT CAAAGCTTTT TAATAGAGAA ACTAACATGT TTGGAGGGGA TTC ATG Met	116
GCC CAA CAT TTA TGG ATT TTG TTG GGA AGT CTC AGT TGC CGA ACA AGC Ala Gln His Leu Trp Ile Leu Leu Gly Ser Leu Ser Cys Arg Thr Ser -15 -10 -5	164
AAC CGG CGG Asn Arg Arg 1	173
(2) INFORMATION FOR SEQ ID NO: 92:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 242 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR 	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 66149 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.1</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 92:	

ACACTTGART TGGGGTTAAG TTGAAGAACA GACAAACTTA GACACAAAGC TATGCAAAAA			
TTGTG ATG AAC AAG GAA RAA GTA AGT TTN GAA AGG ARA GCA CAG GTC AGA Met Asn Lys Glu Xaa Val Ser Xaa Glu Arg Xaa Ala Gln Val Arg -25 -20 -15	110		
TTA TAT TTA TTC TCA GGA TTT TGG ACT TTT KTA TTA GGG AAA TTT AAA Leu Tyr Leu Phe Ser Gly Phe Trp Thr Phe Xaa Leu Gly Lys Phe Lys -10 -5	158		
CAA GGG GAA TGR TCT TAT ATK KGT ATT CTA GAA AGA TTA CTG TGG CAG Gln Gly Glu Xaa Ser Tyr Xaa Xaa Ile Leu Glu Arg Leu Leu Trp Gln 5 10 15	206		
CAG CAG TAT GWA GGA TGG CTT GTA GGR GAT AAG AGA Gln Gln Tyr Xaa Gly Trp Leu Val Gly Asp Lys Arg 20 25 30	242		
(2) INFORMATION FOR SEQ ID NO: 93:			
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 439 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR 			
(ii) MOLECULE TYPE: CDNA			
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate			
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 200361 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6</pre>			
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 93:			
ATTGAAAGAT GGTAAAATGG TGCAGAAGGG GACTTACACT GAGTTCCTAA AATCTGGTAT	60		
AGATTTTGGC TCCCTTTTAA AGAAGGATAA TGAGGAAAGT GAACAACCTC CAGTTCCAGG	120		
AACTCCCACA MYAAGGGAAT CGTACCCTTC TCAGAGTCTT CGGTTTGGTC TCAACAATCT	180		
TCTAGACCCT CCTTGAAAG ATG GTG CTC TGG AGA GCC AAG ATA CAN MGG AAT Met Val Leu Trp Arg Ala Lys Ile Xaa Arg Asn -50 -45	232		
GTC CCA GTT ACA CTA TCA GAG GAG AAC CGT TCT GAA GGA AAA GTT GGT Val Pro Val Thr Leu Ser Glu Glu Asn Arg Ser Glu Gly Lys Val Gly -40 -35 -30	290		
TTT CAG GCC TAT AAG AAT TAC TTC AGA GCT GGT GCT CAC TGG ATT GTC Phe Gln Ala Tyr Lys Asn Tyr Phe Arg Ala Gly Ala His Trp Ile Val -25 -20 -15	328		

71	
TTC ATT TTC CTT ATT CTC CTA AAC ACT GCA GCT CAG GTT GCC TAT GTG Phe Ile Phe Leu Ile Leu Leu Asn Thr Ala Ala Gln Val Ala Tyr Val -10 5	376
CTT CAA GAT TGG TGG CTT TCA TAC TGG GCA AAC AAA CAA AGT ATG CTA Leu Gln Asp Trp Trp Leu Ser Tyr Trp Ala Asn Lys Gln Ser Met Leu 10 15 20	424
AAT GTC ACT GTA AAT Asn Val Thr Val Asn 25	439
(2) INFORMATION FOR SEQ ID NO: 94:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 232 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: DOUBLE(D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 125178 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 94:	
ATGTAGTGAA TAAAGTTTGA GAACCACTGA CTTGAACTTT AGCATGATTT GATACACAGG	60
GTCCTCTGTA ATCGTACTTC GTTCTGCTTT AAGGCTGTTG GGCTGTCTCC TCCAACCCAT	120
CCKK ATG TTG TAK TTT TTC ACC TCK GTC CTT TGG CTT ACG TCA CCN Met Leu Leu Xaa Phe Phe Thr Ser Val Leu Trp Leu Thr Ser Pro -15 -10 -5	169
TCC CAA CCT AAT ACC TGC CCT TCT AGT CTT CTG TGT ACT TAT CCA AAT Ser Gln Pro Asn Thr Cys Pro Ser Ser Leu Leu Cys Thr Tyr Pro Asn 1 5 10	217

232

(2) INFORMATION FOR SEQ ID NO: 95:

CTA AAC CCT CCA TGG

Leu Asn Pro Pro Trp

15

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 229 base pairs

7.	2
(B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
DLECULE TYPE: CDNA	

- (ii) MC
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 140..205
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq IILGCLALFLLLQ/RK

229

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 95:

AACAGTTACG AAGGAGAGCT GCAAAAGTTG CAGCAGAAAG GTTGGGAGTC CCGACAGGTT 60 CCGTAGCCCA CAGAAAAGAA GCAAGGGACG GCAGGACTGT TTCACACTTT TCTGCTTCTG 120 GAAGGTGCTG GACAAAAAC ATG GAA CTA ATT TCC CCA ACA GTG ATT ATA ATC Met Glu Leu Ile Ser Pro Thr Val Ile Ile Ile -20 CTG GGT TGC CTT GCT CTG TTC TTA CTC CTT CAG CGG AAG AAT TTG CGC Leu Gly Cys Leu Ala Leu Phe Leu Leu Leu Gln Arg Lys Asn Leu Arg 220 -10 AGA CCC TGG

(2) INFORMATION FOR SEQ ID NO: 96:

Arg Pro Trp

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 292 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 134..274
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq TWLGLLSFQNLHC/FP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 96:

ATCATTTTCT TATCCCTGCT GATTTCAAAC CTTCCCATGG TTTAGAAGCA TAACCTGTAA	60
TGTAATGCAA GTCCCCTAAC TCCCTGGTTG CTAACATTAA CTTCCTTAAG TAATAATCAA	120
TGAAAGAVAT TCT ATG CAT GGT TTT GAA ATA ATA TCC TTG AAA GAG GAA Met His Gly Phe Glu Ile Ile Ser Leu Lys Glu Glu -45 -40	169
TCA CCA TTA GGA AAG GTG AGT CAG GGT CCT TTG TTT AAT GTG ACT AGT Ser Pro Leu Gly Lys Val Ser Gln Gly Pro Leu Phe Asn Val Thr Ser -35 -20	217
GGC TCA TCA TCA CCA GTG ACC TGG TTG GGC CTA CTC TCC TTC CAG AAC Gly Ser Ser Pro Val Thr Trp Leu Gly Leu Leu Ser Phe Gln Asn -15 -10 -5	265
CTG CAT TGC TTC CCA GAC CTC CCC GGG Leu His Cys Phe Pro Asp Leu Pro Gly 1 5	292
(2) INFORMATION FOR SEQ ID NO: 97:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 458 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Cancerous prostate (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 270437 (C) IDENTIFICATION METHOD: Von Heijne matrix	
(D) OTHER INFORMATION: score 5.9 seq NTLFLHLSGLSAA/DT	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 97:	
AAGCTCTGAG ACAGGAGCCC AGCCCTGGGA TTTTCAGGTG TTTTCATTTG GTGGTCAGGC	60
CTGAACAGAG TGTTTTCCTT TGGTGGTCAG GACTGAGCAG AGAGACCTCA CCATGGAGCT	120
KGGSYGKTG CKGGCTTTTT CTTGTGGCCA TTTTGAAAGA TGTCCGGTCT GAGGGACAAC	180
PATTGGAATC TGGGGGAAGT TCGGTCCAGC CCGGGGAGTC CCTGCGACTC TCCTGTGCAG	240
CCGCTGGATT CGCNTTTCGC AATTTTGCC ATG ACT TGG GTC CGC CAC GCT CCA Met Thr Trp Val Arg His Ala Pro -55 -50	293
GGG AAG AGT CTG GAA TGG GTC GCA ACC GTC ACA GAT GGT GGT GAT AAG Gly Lys Ser Leu Glu Trp Val Ala Thr Val Thr Asp Gly Gly Asp Lys	341

ACC TTT TAT GCG GCC TCC GTG AAG GGC CGC TTC AAC GTC TCC AGG GAC Thr Phe Tyr Ala Ala Ser Val Lys Gly Arg Phe Asn Val Ser Arg Asp -30 -25 -20	389											
AAT TCC AAG AAC ACG TTA TTT CTG CAT TTG AGC GGC CTG AGT GCC GCC Asn Ser Lys Asn Thr Leu Phe Leu His Leu Ser Gly Leu Ser Ala Ala -15 -5	437											
GAC ACG GGC TGG TGG GGG ATC Asp Thr Gly Trp Gly Ile 1 5	458											
(2) INFORMATION FOR SEQ ID NO: 98: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 226 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Prostate (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 143184 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: SCORE 5.8 seq LTSFFSLTANCQS/AG (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 98:												
AACATACCCT TCAGGTTTAG GTCTTTCTTA GGTAAAGTTT TAACTTTAGT ATATCTTCCT	60											
CAGGGCGGCC TTCTCCTTCC CCCTAGTAAG TGRAGAAACC CTTGTGTKTC TGCCCTCTGA	120											
ACTCACCGCA TTTGGGATTA CC ATG CTA ACA TCC TTT TTT TCA CTG ACT GCA Met Leu Thr Ser Phe Phe Ser Leu Thr Ala -10 -5	172											
AAT TGC CAG AGT GCA GGA ACT ATC TCA TTT GCT GCT TTC TCC CTA ATG Asn Cys Gln Ser Ala Gly Thr Ile Ser Phe Ala Ala Phe Ser Leu Met 1 5 10	220											
CCT GGA Pro Gly	226											
(2) INFORMATION FOR SEQ ID NO: 99:												

- - (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 140 base pairs
 (B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Cancerous prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 72125 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.8</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 99:	
ACTTCCCTTC CCCCTCTAGC ATTGCTACCT TCTCTCCTAC ACGCACGCAG GCATATAAAC	60
GTAGGTTTTT G ATG CTC CTC TGC CTG TTG ACC CCG CTA TTT TTC ATG TTK Met Leu Leu Cys Leu Leu Thr Pro Leu Phe Phe Met Xaa -15 -10	110
CCA ACA GGT TTT TCT TCC CCC AGT CCT GGG Pro Thr Gly Phe Ser Ser Pro Ser Pro Gly -5 1 5	140
(2) INFORMATION FOR SEQ ID NO: 100:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 288 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Cancerous prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 178240 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.7</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 100:	
AATTGGCGCG GGGCGTCCGT AGCCACGGCA ACAGGTTGCT TCTGCAGTCT GAGCTGAGCG	60
CCTTTCGCAC GACTTGGAGT TACGGTTTAT TTGATACCCC GGTACCCCTA CGCAAGCAAG	120
CCCACATCGA CACACATTCA CACACGCCCT TCAGCACCC CTCCCAGCAC CACGACC	177

ATG Met	GAC Asp -20	GAC Asp	GAC Asp	TAT Tyr	GAA Glu	GCG Ala -15	TAC Tyr	CAC His	AGT Ser	CTG Leu	TTC Phe -10	TTG Leu	TCG Ser	CTG Leu	CTC Leu	225
GGA Gly -5	CTC Leu	TGC Cys	CCG Pro	TCT Ser	AAG Lys l	ACT Thr	CCC Pro	ATC Ile	AAT Asn 5	GAA Glu	AAT Asn	GCT Ala	CCC Pro	GTC Val 10	TTT Phe	273
			CCG Pro 15													288

(2) INFORMATION FOR SEQ ID NO: 101:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 393 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 298..354
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.7

seq WLVWLLLGHMVVS/QM

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 101:

CTCTTGCCTC	AGGCTTGGAG GCCTCCGAGC AGCAACATCG TCCCAATTAT ACCCCGTTGG	60
AGCATCTTCA	GATCTTCCAC TCTTTTCACA ACGCAATCAA AATCTTCGTA CCCATTTTGC 1	20
AGTAGTGATC	TCTAAACTCT CAGCGTAGGC ATCGGGAACC TTCGTGCCAA GGAGCCATGC 1	80
TGCCCCGATG	GGAACTGGCA CTTTACCTAC TTGCCTCACT AGGCTTCCAC TTCTATTCCT 2	240
TCTATTAAGT	TTACAAAGTC TCCAGAGGAT GCGACCGACT TTGAGTGGAG CTTCTGG 2	97
ATG GAA TGG Met Glu Trp	G GGG AAG CAG TGG CTG GTG TGG CTT CTC CTT GGC CAC ATG O Gly Lys Gln Trp Leu Val Trp Leu Leu Gly His Met -15 -10 -5	345
GTA GTG TCT Val Val Ser	CAA ATG GCC ACA CTG CTG GCA AGA AAG CAC AGA CCC TGG Gln Met Ala Thr Leu Leu Ala Arg Lys His Arg Pro Trp 1 5 10	193

- (2) INFORMATION FOR SEQ ID NO: 102:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 281 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: DOUBLE(D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 135251 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.7</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 102:	
ATATACAGAG AATAAACGTC ATCCCTCTAA CATTAATATG TTCAGTTTTA TGTACCTGAG	60
AGTTGATGGT TTAATTTGTG GGTTTGCCCA GACTCTCTTG CGACTTCTCT CATCATCTGC	120
FCTTTAGCAC TTCC ATG AGA CGG GGC AAG AGA TTG TTG GAG TCT CAA TCC Met Arg Arg Gly Lys Arg Leu Leu Glu Ser Gln Ser -35 -30	170
AGC AGC CCG AAA GCC TGT CTG CAG CTT GGG TTT GAG ACT GAA CTA ACT Ser Ser Pro Lys Ala Cys Leu Gln Leu Gly Phe Glu Thr Glu Leu Thr -25 -20 -15	218
CAG GGT GTT TTG TGG ATT TTA GTT ATC CAG GCT GTC CCT GTT CCC TCA Sin Gly Val Leu Trp Ile Leu Val Ile Gin Ala Val Pro Val Pro Ser -10 -5 1 5	266
TTA ACA AAA ACA AAA Leu Thr Lys Thr Lys 10	281
-	
(2) INFORMATION FOR SEQ ID NO: 103:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 276 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 205264 (C) IDENTIFICATION METHOD: Von Hoijno matrix</pre>	

WO 99/06550 PCT/IB98/01232

(D) OTHER INFORMATION: score 5.7 seq ALLESVVWLPCHG/RG	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 103:	
AGACCAGGCC CATTTCTCAG AAGCCTTTGG CTCCCCTGAG ATGCCAAATA GCCGCTCACT	60
CTTCCGCCTC CACGGACTGG CTTTGGTGTT CATGCTGGTT GGGATGTCTA CTATGGACCT	120
GCTGAGCACA GGGCTGGGTT CCTGGGGCAC AGAGTTGATG CTTATGGCCC AGGAACTGCT	180
GGGCCCCAGG ACTGGGCGGT TTCC ATG GTT GCT GCC ACA GAA GCA GCA TTG Met Val Ala Ala Thr Glu Ala Ala Leu -20 -15	231
CTG GAG TCA GTA GTG TGG CTG CCT TGC CAT GGC CGT GGT GGG TCT Leu Glu Ser Val Val Trp Leu Pro Cys His Gly Arg Gly Gly Ser -10 -5 1	276
(2) INFORMATION FOR SEQ ID NO: 104: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 421 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 356412 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.6 seq VSLPLLSSWGSTA/WT (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 104:	
AATTACAGCT CTACAATGCA CCAGACGGAC CCATCTGGAT TCTTTCGGGG CTCTTAGCCC	60
	120
AGTTGATGCA CGGCATGGCA CAGGATGCTG GGCAAGGCTG GCAGGCCCGG GAGAGCCTGT	180
·	240
TCCTAACTTC CTCACGGCAC ACCATTCTGC CGTCCTGAGT CTTCTCAAGG TTGGAAGGTG	300
CCCAGATCCA GGGAGATGGT GCTGGCTCTT TGGTGGCTGT GGAGTGTCCA GACAG ATG Met	358
AGC TGG AAT CCT TCA GTT TCT CTG CCT CTC CTG TCA AGT TGG GGT AGC Ser Trp Asn Pro Ser Val Ser Leu Pro Leu Leu Ser Ser Trp Gly Ser	406

,	WO 9	9/0655	50								PCT/IB98/01232					
			-15					-10		79			-5			
			ACT Thr										Ī			421
(2)	INF	ORMA	TION	FOR	SEQ	ID:	NO:	105:								
	(:	i) S	(B) (C)	NCE (LEN(TYPI STRA TOP(STH: E: NU ANDE	193 JCLE: NES	base IC A	e pa: CID DUBL								
	(:	ii) l	MOLE	CULE	TYPE	E: CI	ONA									
	(-	vi) (NAL ORGA	NISI	1: Ho										
	(:	ix) l	(B) (C)	NAME LOCA I DEN	TION TIFI	1: 53 CAT	31 [ON N	l 8 1Etho	DD: <i>\</i>	/on f ce 5. LILI	6					
	()	ki) S	EQUE	ENCE	DESC	RIP	rion	: SE	Q ID	NO:	105	:				
ACA	AATA	raa (CTAAT	rgag <i>i</i>	AT TA	AAA?	ATTI	AA A	CAGG'	IGTC	TGA	ГААТ	CCT :		IG AAG et Lys	58
AGA Arg -20	ATT Ile	CAG Gln	GGG Gly	ATA Ile	TTG Leu -15	TTC Phe	CTG Leu	ATT Ile	TTG Leu	CTT Leu -10	TCT Ser	CTC Leu	CAC His	TTG Leu	GAA Glu -5	106
AGG A rg	AGG Arg	TGG Trp	ACG Thr	AGC Ser 1	CCA Pro	TCA Ser	GAC Asp	CAC His 5	AGC Ser	CTG Leu	TTG Leu	CTA Leu	GGA Gly 10	GGA Gly	AAT Asn	154
TCC Ser	TTG Leu	GCT Ala 15	CAA Gln	CAT His	GCA Ala	GAA Glu	AGT Ser 20	GTA Val	GTA Val	CGC Arg	CAA Gln	GGG Gly 25				193
(2)	INFO	ORMA?	rion	FOR	SEQ	ID 1	NO: 3	106:								
	į)	L) SE	(B)	ICE (LENG TYPE STRA	TH: :: NU	435 CLE	base IC AC	e pai								

(D) TOPOLOGY: LINEAR

(A) ORGANISM: Homo Sapiens

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 298..402
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 5.5

seq LLTFGLEVCLAAG/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 106:

AAAGGAAGGG GGGGCGGAAC CAGCCTGCAC GCGCTGGCTC CGGGTGACAG CCGCGCGCCT	60
CGGCCAGGAT CTGAGTGATG AGACGTGTCC CCACTGAGGT GCCCCACAGC AGCAGGTGTT	120
GAGCATGGGC TGAGAAGCTG GACCGGCACC AAAGGGCTGG CAGAAATDVG CGCCTGGCTG	180
ATTCCTAGGC AGTTGGCRGC AGCAAGGAGG AGAGGCCGCA GCTTCTGGAG CAGAGCCGAG	240
ACGAAGCAGT TCTGGAGTGC CTGAACGGCC CCCTGAGCCC TACCCGCCTG GCCCACT	297
ATG GTC CAG AGG CTG TGG GTG AGC CGC CTG CTG CGG CAC CGG AAA GCC Met Val Gln Arg Leu Trp Val Ser Arg Leu Leu Arg His Arg Lys Ala -35	345
CAG CTC KKG CTG GKC AAC CTG CTA ACC TTT GGC CTG GAG GTG TGT TTG Gln Leu Xaa Leu Xaa Asn Leu Leu Thr Phe Gly Leu Glu Val Cys Leu -15 -5	393
GCC GCA GGA TCA CCT ATG TGC CGC CTC TGC TGC TGG AAG TGG Ala Ala Gly Ser Pro Met Cys Arg Leu Cys Cys Trp Lys Trp 1 5 10	435

(2) INFORMATION FOR SEQ ID NO: 107:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 392 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 27..80
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.5

seq PFALVTSCSSVFS/GD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 107:

01

	Met Ala Ala Gly Val Pro Phe Ala Leu -15 -10															
					TCC Ser											101
					GAT Asp											149
					TGG Trp											197
					GTG Val 45											245
					GCA Ala											293
					GTC Val											341
					ATC Ile											389
ATA Ile																392

- (2) INFORMATION FOR SEQ ID NO: 108:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 358 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 290..331
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.5

seq TVFLXFCFPRCHS/DS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 108:

TCAAGTTTTA ACGAAGAAAA ACATCATTGC AGTGAAATAA AAAATTTTAA AATTTTAGAA	120
CAAAGCTAAC AAATGGCTAG TTTTCTATGN TTCTTCTTCA AACGCTTTCT TTGAGGGRGM	180
AAGAGTCAMA CAAACAAGCA GTTTTACCTA AAATAAAGAA CTAGTTTTAG AGGTCAGAMG	240
AMAGGMGCAA GTTTTGCGAG WGGCACGGAA GGAGTGTGCT GGCAGTACA ATG ACA GTT Met Thr Val	298
TTC CTT TMN TTT TGC TTT CCT CGC TGC CAT TCT GAC TCA CAT ARG RTG Phe Leu Xaa Phe Cys Phe Pro Arg Cys His Ser Asp Ser His Xaa Xaa -10 5	346
CAG CAA TCA GCG Gln Gln Ser Ala	358

(2) INFORMATION FOR SEQ ID NO: 109:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 310 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 44..187
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.4

seq ILLEVFVWNGLQG/LP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 109:

AAS'	TTCT'	rcc '	TGCC	AAGA(GA A	CAAT	GCCG	A GA	AACA	GAGC	GAA		К ТТ Хаа			55
AAT Asn	TTT Phe	TGG Trp	CAA Gln	AAA Lys -40	CTT Leu	GGA Gly	AGA Arg	AAA Lys	AAA Lys -35	CCC Pro	CGC Arg	ATA Ile	TTT Phe	ACC Thr -30	TGT Cys	103
ACC Thr	CAG Gln	AGC Ser	TCC Ser -25	ACA Thr	GGT Gly	GAG Glu	GCG Ala	GCA Ala -20	GTT Val	AAA Lys	GCA Ala	GAA Glu	AAT Asn -15	CTA Leu	ATT Ile	151
CTT Leu	CTG Leu	GAA Glu -10	GTT Val	TTT Phe	GTC Val	TGG Trp	AAC Asn -5	GGA Gly	CTC Leu	CAG Gln	GGT Gly	CTT Leu 1	CCT Pro	TCG Ser	GAG Glu	199
CTG Leu	TCA Ser	GAT Asp	ACA Thr	AGT Ser	GGA Gly	TCC Ser	TCT Ser	AAG Lys	AAA Lys	CTT Leu	GGG Gly	AGC Ser	CTT Leu	GTG Val	GGC Gly	247

WO 99/06550		83	PCT/IB98/01232
5	10	15	20
TGG TGG AGA ACT CTC Trp Trp Arg Thr Let 25	ı Lys Met Ala	CCA GCC TGT CTA TGG TG Pro Ala Cys Leu Trp Se 30	CT ATG TGG 295 er Met Trp 35
GAA TCA CCG CCA CGC Glu Ser Pro Pro Arc 40			310
(2) INFORMATION FOR	R SEQ ID NO: 1	10:	
(A) LEN (B) TYP (C) STR	CHARACTERISTIC GTH: 284 base E: NUCLEIC ACT ANDEDNESS: DOC OLOGY: LINEAR TYPE: CDNA	pairs ID	
	ANISM: Homo Sa	apiens cerous prostate	
(B) LOC (C) IDE (D) OTH	E/KEY: sig_per ATION: 66173 NTIFICATION ME ER INFORMATION	3 ETHOD: Von Heijne matr	
AAGTCCAGAG GCCTGGCC	CT GCCAAGAAGG	CGCTCTCCGG AATCAACACC	TGGGGGCTTG 60
GAAGG ATG TTT CGC T Met Phe Arg S -35	CA GAT CGA ATO er Asp Arg Met -30	G TGG ARC TGC CAT TGG t Trp Xaa Cys His Trp 0 -25	AAA TGG AAG 110 Lys Trp Lys
CCC AGT CCT CTC CTC Pro Ser Pro Leu Leu -20	TTC TTA TTT (Phe Leu Phe A	GCT TTA TAT ATC ATG TG Ala Leu Tyr Ile Met Cy -10	T GTT CCT 158 s Val Pro
CAC TCA GTG TGG GGA His Ser Val Trp Gly -5	TGT GCC AAC 1 Cys Ala Asn (IGC CGA GTG GTT TTG TC Cys Arg Val Val Leu Se 5	C AAC CCT 206 r Asn Pro 10
TCT GGG ACC TTT ACT Ser Gly Thr Phe Thr 15	TCT CCA TGC TS Ser Pro Cys T	TAC CCT AAC GAC TAC CC Tyr Pro Asn Asp Tyr Pr 20	A AAC AGC 254 o Asn Ser 5
CAG GCT TGC ATG TGG Gln Ala Cys Met Trp 30			284

		(i) :	(A) (B) (C)	LEN TYP STP	NGTH:	: 398 NUCLE EDNES	B bas EIC A ES: [OOUBI	airs							
		(ii)	MOLE	ECULE	E TYE	PE: (DNA									
		(vi)	(A)	ORG	SANIS	M: H	omo	Sapi ormal	ens pro	stat	e					
	<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 123215 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.3</pre>															
	(xi)	SEQU	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	111	:				
TCC	TTCA	TCT	TGTG	ттст	AA A	ACCT	TGCA	A GT	TCAG	GAAG	AAA	CCAT	CTG	CATC	САТАТТ	60
															TCTACA	120
AC	Met	ACC Thr -30	CAA Gln	AGG Arg	AGC Ser	Ile	GCA Ala -25	GGT Gly	CCT Pro	ATT Ile	Cys .	AAC Asn -20	CTG Leu	AAG :	TTT Phe	167
GTG Val	ACT Thr -15	reu	CTG Leu	GTT Val	GCC Ala	TTA Leu -10	AGT Ser	TCA Ser	GAA Glu	CTC Leu	CCA Pro -5	TTC Phe	CTG Leu	GGA Gly	GCT Ala	215
GGA Gly 1	GTA Val	CAG Gln	CTT Leu	CAA Gln 5	GAC Asp	AAT Asn	GGG Gly	TAT Tyr	AAT Asn 10	GGA Gly	TTG Leu	CTC Leu	ATT Ile	GCA Ala 15	ATT Ile	263
AAT Asn	CCT	CAG Gln	GTA Val 20	CCT Pro	GAG Glu	AAT Asn	CAG Gln	AAC Asn 25	CTC Leu	ATC Ile	TCA Ser	AAC Asn	ATT Ile 30	AAG Lys	GAA Glu	311
ATG Met	ATA Ile	ACT Thr 35	GAA Glu	GCT Ala	TCA Ser	TTT Phe	TAC Tyr 40	CTA Leu	TTT Phe	AAT Asn	GCT Ala	ACC Thr 45	AAG Lys	AGA Arg	AGA Arg	359
GTA Val	TTT Phe 50	TTC Phe	AGA Arg	AAT Asn	ATA Ile	AAG Lys 55	ATT Ile	TTA Leu	ATA Ile	CCT Pro	GCC Ala 60	CAG Gln				398
(2)	INFO	ORMAT	rion	FOR	SEQ	ID N	1 0: 1	112:								
	i)	.) SE	(A) (B) (C)	LENG TYPE STRA	TH:	324 CLEI NESS	base C AC : DC	pai ID UBLE								

WO 99/06550 PCT/IB98/01232

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens
(F) T\(\frac{1}{2}\)SUE TYPE: Prostate

(ix) FEATURE:

(A) NAME/KEY: sig_peptide
(B) LOCATION: 187..228

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.3

seq IIPLLLLRSACN/VH

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 112:

ACTCCAGGAG CCGGGACCAA AATAACCGGG CGGGAGGGGA CACCTCGCAG AGATGGATCT 60

CGAACTCCTG GGCTCAAGCG ATCCTTTCAC CTTGGCCTCT CAAGTAGCTG GGACCACATT 120

TGCTCACCAG CTGGCCCAAG ACCAGACTGG GCAACATGGG TCATCCTCCT CTAAGATTCC 180

AGGACC ATG ATC ATC CCT CTA TTG CTA CTT CTT AGA TCA GCT TGT AAT

Met Ile Ile Pro Leu Leu Leu Leu Arg Ser Ala Cys Asn

-10

-5

GTC CAT CTC CCC CAC CAG ACT GCG TCT CCA GCA TCT CTG AGT CCC CAG

Val His Leu Pro His Gln Thr Ala Ser Pro Ala Ser Leu Ser Pro Gln

1 5 10 15

GGC CTG GCC TGG GGC TTG CTA CAT GGT GGG TGC TCA GTA ACT GTG AGA
Gly Leu Ala Trp Gly Leu Leu His Gly Gly Cys Ser Val Thr Val Arg
20 25 30

(2) INFORMATION FOR SEQ ID NO: 113:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 293 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 231..287
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.3

seq VLLLSXNLNLIIQ/SS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 113:

•	
TTGGAGCAAG TGAGAAGACA AGTKAGAGGT AAGCWGKTRT TGAGAATAGG GGKCTGATTG	120
TGCCAGCTTT GTATACVATT ATNAGGAACN DGGACTTTGT CCTGAAGGTA ACTGGGCAAT	180
TGTTGAGGTC ACCACCATCT ACTGTCTGGA TTACCGAGGA AACTTTCTAA ATG TMS Met Xaa	236
TCT CCA CTT CCA GTC CTG CTC CTC TCA TKC AAT CTC AAC CTA ATA ATT Ser Pro Leu Pro Val Leu Leu Ser Xaa Asn Leu Asn Leu Ile Ile -15	284
CAG AGT AGT Gln Ser Ser 1	293
(2) INFORMATION FOR SEQ ID NO: 114:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 402 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 244381 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.2 seq LLTFLVFTXKLSS/LN</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 114:	
ACACTGAAAT CAATCTGTTC AATAGCATTA TACCATATTT GACATACCAT AGCCATGTTA	60
ATCTGATATT GTAGAATAGC ATAGTAKAAT AATAATAACT CCTAACTCAA GGATGTTGWG	120
WKCCTTTATA ACCAGCAATC CATGTTARAT ATTAGCACAG TGCCTAAAAC ATATTAAGCA	180
TTCAATAAAT GATCGCTACT ATTTTTACTA ACATCCTACA GATTTGGAAA TTGAGTCTTA	240
GAA ATG TTA ATG TGT AAA ATG CTA AAG AGC CAA AAA AAC TGC CAG GAA Met Leu Met Cys Lys Met Leu Lys Ser Gln Lys Asn Cys Gln Glu -45 -35	288
AAT ATR ARA ATT AAA ATC ATT TTA TTT CTG AAA CCC ATG TGT TCC CCC Asn Xaa Xaa Ile Lys Ile Ile Leu Phe Leu Lys Pro Met Cys Ser Pro -30 -25 -20	336
CAA TAT CTT CTA ACA TTT CTA GTA TTT ACA GRA AAA CTT TCA AGT CTC Gln Tyr Leu Leu Thr Phe Leu Val Phe Thr Xaa Lys Leu Ser Ser Leu -15	384

ΑT	r A	ATC	RGA	AAG	TTT	CAT

(2) INFORMATION FOR SEO ID NO: 115:

Asn Ile Xaa Lys Phe His

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 470 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 306..461
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.2

seq IIVILHCAASIIS/CP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 115:

CTA TTA CTT GCC ATT GAT CTT AAA ATT ATA GTT ATC CTC CAC TGT GCT
Leu Leu Leu Ala Ile Asp Leu Lys Ile Ile Val Ile Leu His Cys Ala
-20 -15 -10

GCA TCC ATA ATT TCA TGT CCC TCA
Ala Ser Ile Ile Ser Cys Pro Ser
-5

(2) INFORMATION FOR SEQ ID NO: 116:

- (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 334 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 116..184
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.1

seq ATSVSLEAQSCFA/WP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 116:

ATTTTTGAAA ACTGTAATGC TTTAAAACTT ACTTTATTGG ATCTCTTTGC AGCTTTTGAC	60
ACAGTGAACC ACTTTCCTTT CCTGAAATGC TTTCCTCTCT TGGCTTTCTG ATGCC ATG Met	118
TTC TCC TGT TTC TCT ACT TCT CTG GCC ACT TCT GTC TCC TTA GAA Phe Ser Cys Phe Phe Ser Thr Ser Leu Ala Thr Ser Val Ser Leu Glu -20 -15 -10	166
GCT CAG TCT TGC TTT GCC TGG CCC TTG ATT GTT AGT TTT CCC CAG GGC Ala Gln Ser Cys Phe Ala Trp Pro Leu Ile Val Ser Phe Pro Gln Gly -5	214
TCA CTT CTT AGC CCC TTT CTC CTC ATG TCT TAT AAT TTG AGT CAT CTC Ser Leu Leu Ser Pro Phe Leu Leu Met Ser Tyr Asn Leu Ser His Leu 15 20 25	262
ATC TAC TCT GGG GAG TTG AAT GGT CGC TTG TAT GCT GAA AAC TCC CAA Ile Tyr Ser Gly Glu Leu Asn Gly Arg Leu Tyr Ala Glu Asn Ser Gln 30 35 40	310
ATT TGT ATC TGT AGC CCA GCC GGG Ile Cys Ile Cys Ser Pro Ala Gly	334

(2) INFORMATION FOR SEQ ID NO: 117:

45

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 302 base pairs

50

- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 78..227
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.1

seq RTALILAVCCGSA/SI

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 117:

AGTTTCCAAG GGAAGGAGCA GCGTGTGGGA AAGCACAGAA GAGTGAGAAG GAAGCGACTA 60

AATTTTATTT ACTTTCT ATG CAT CAT GGC CTC ACA CCA CTG TTA CTT GGT

Met His His Gly Leu Thr Pro Leu Leu Gly

-50

-45

GTA CAT GAG CAA AAA CAG CAA GTG GTG AAA TTT TTA ATC AAG AAA AAA 158
Val His Glu Gln Lys Gln Gln Val Val Lys Phe Leu Ile Lys Lys Lys
-35 -30 -25

GCA AAT TTA AAT GCA CTG GAT AGA TAT GGA AGA ACT GCT CTC ATA CTT

Ala Asn Leu Asn Ala Leu Asp Arg Tyr Gly Arg Thr Ala Leu Ile Leu

-20

-15

-10

GCT GTA TGT TGT GGA TCG GCA AGT ATA GTC AGC CTT CTA CTT GAG CAA 254
Ala Val Cys Cys Gly Ser Ala Ser Ile Val Ser Leu Leu Glu Gln
-5 1 5

AAC ATT GAT GTA TCT TCT CAA GAT CTA TCT GGA CAG ACG GCC CCC GGG
Asn Ile Asp Val Ser Ser Gln Asp Leu Ser Gly Gln Thr Ala Pro Gly
10 25

- (2) INFORMATION FOR SEQ ID NO: 118:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 381 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 319..369
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.1

seq IYFFACFQALTSS/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 118:

AAGACTGGAC AAAGGGGGTC ACACATTCCT TCCATACGGT TGAGCCTCTA CCTGCCTGGT	120
GCTGGTCACA GTTCAGCTTC TTCATGATGG TGGATCCCAA TGGCAATGAA TCCAGTGCTA	180
CATACTTCAT CCTAATAGGC CTCCCTGGTT TAGAAGAGGC TCAGTTCTGG TTGGCCTTCC	240
CATTGTGCTC CCTCTACCTT ATTGCTGTGC TAGGTAACTT GACAATCATC TACATTGTGC	300
GGACTGAGCA CAGCCTGC ATG AGC CCA TGT ATA TAT TTC TTT GCA TGC TTT Met Ser Pro Cys Ile Tyr Phe Phe Ala Cys Phe -15 -10	351
CAG GCA TTG ACA TCC TCA TCT CCA CCT CAG Gln Ala Leu Thr Ser Ser Ser Pro Pro Gln -5	381
(2) INFORMATION FOR SEQ ID NO: 119: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 318 base pairs	
(B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Hypertrophic prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 49141 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.1</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 119:	
CTTTCTGTGT CTCCTTTCCT CCGCCTCAGT TTGGGGCGGG TCGGGGGA ATG GCT GAG Met Ala Glu -30	57
GAG ATG GAG TCG TCG CTC GAG GCA AGS TTT TCG TCC AGC GGG GCA GTG Glu Met Glu Ser Ser Leu Glu Ala Xaa Phe Ser Ser Gly Ala Val -25 -20 -15	105
TCA GGG GCC TCA GGG TTT TTG CCT CCT GCC CGC TCC CGC ATC TTC AAG Ser Gly Ala Ser Gly Phe Leu Pro Pro Ala Arg Ser Arg Ile Phe Lys -10 -5 1	153
ATA ATC GTG ATC GGC GAC VBC AAT GTG GGC AAG ACA TGC CTG ACC TAC Ile Ile Val Ile Gly Asp Xaa Asn Val Gly Lys Thr Cys Leu Thr Tyr 10 15 20	201
CGC TTC TGC GCT GGC CGC TTC CCC GAC CGC ACC GAG GCC ACG ATA GGG Arg Phe Cys Ala Gly Arg Phe Pro Asp Arg Thr Glu Ala Thr Ile Gly	249

WO 99/06550		91	PCT/IB98/01232					
25		30	35					
GTG GAT TTC CGA GAA Val Asp Phe Arg Glu 40	CGA GCG GTG GAG Arg Ala Val Glu 45	Ile Asp Gly Glu	CGC ATC AAG 297 Arg Ile Lys 50					
ATC CAG CTA TGG GAC Ile Gln Leu Trp Asp 55			318					
(2) INFORMATION FOR	SEQ ID NO: 120:							
(A) LENG (B) TYPI (C) STR	CHARACTERISTICS: STH: 243 base pa E: NUCLEIC ACID ANDEDNESS: DOUBL DLOGY: LINEAR							
(ii) MOLECULE	TYPE: CDNA							
	SOURCE: ANISM: Homo Sapi SUE TYPE: Hypert							
(ix) FEATURE:								
 (A) NAME/KEY: sig_peptide (B) LOCATION: 61153 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.1 seq VSGASGFLPPARS/RI 								
(xi) SEQUENCE	DESCRIPTION: SE	Q ID NO: 120:						
AAATCTCTCA GCCTTTCT	GT GTCTCCTTTC CT	CCGCCTCA GTTTGGGG	CG GGTCGGGGGA 60					
ATG GCT GAG GAG ATG Met Ala Glu Glu Met -30								
GGG GCA GTG TCA GGG Gly Ala Val Ser Gly -15								

(2) INFORMATION FOR SEQ ID NO: 121:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 278 base pairs

(B) TYPE: NUCLEIC ACID

	(C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii)	MOLECULE TYPE: CDNA	
(vi)	ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate	
(ix)	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 153233 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5 seq HLSLILLKPLCLP/NN	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 121:	
ACCTTTTATA	AACATTTTGT TTAACTTTTA TTGTGGTAAA ATACACATAA CACTTCTCTT	60
CTTTTAGACC	TGGGCTGGTA AGAAGTGCTG AAGATGTTTT TTAGAGATTT GTGGTATGAC	120
AAATTCCACT	GGGGTTTCTG ASCTTCTCAG TC ATG CTT GTC TTG GGG TCA CCA Met Leu Val Leu Gly Ser Pro -25	173
CTC CTT GGC Leu Leu Gly -20	C CCT CTC CTA TGG CAC CTG TCC CTC ATT CTG CTC AAG CCC y Pro Leu Leu Trp His Leu Ser Leu Ile Leu Leu Lys Pro -15 -10 -5	221
CTA TGC CTT Leu Cys Leu	F CCC AAC AAC TTG CCT TTA GCT CTG GGC AGA TGT CTT TGC Pro Asn Asn Leu Pro Leu Ala Leu Gly Arg Cys Leu Cys 1 5 10	269
TTG CAC TCG Leu His Ser 15	•	278
(2) INFORMA	ATION FOR SEQ ID NO: 122:	
(i) S	GEQUENCE CHARACTERISTICS: (A) LENGTH: 301 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii)	MOLECULE TYPE: CDNA	
(vi)	ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Cancerous prostate	
(ix)	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 56220 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5	

seq VLFMTTAVDLVIT/EV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 122:

AGA	AAGG'	IGT '	TTTG(GTCT'	TC T	CCTT	AGTC	C AG	GAAA	AGAT	GTA	CGAA	ATA (GTGA	C ATG Met -55	58
CAC His	TTA Leu	TTA Leu	GAT Asp	TTG Leu -50	GAA Glu	TCT Ser	ATG Met	GGC Gly	AAA Lys -45	AGT Ser	TCA Ser	GAT Asp	GGA Gly	AAG Lys -40	TCG Ser	106
TAT Tyr	GTT Val	ATT Ile	ACG Thr -35	GGG Gly	AGC Ser	TGG Trp	AAT Asn	CCA Pro -30	AAA Lys	TCC Ser	CCA Pro	CAT His	TTT Phe -25	CAA Gln	GTT Val	154
GTA Val	AAT Asn	GAA Glu -20	GAA Glu	ACT Thr	CCT Pro	AAA Lys	GAT Asp -15	AAA Lys	GTC Val	CTG Leu	TTT Phe	ATG Met -10	ACC Thr	ACA Thr	GCT Ala	202
				ATA Ile												250
				CGC Arg 15												298
GCG Ala																301

(2) INFORMATION FOR SEQ ID NO: 123:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 129 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- -(ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 1..63
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8

seq VLFVFSSIPLTFL/FQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 123:

ATG GAG AAT TTG AAA GAC TTT TAT GTG TTG TTT GTA TTC TCT AGC ATT

Met Glu Asn Leu Lys Asp Phe Tyr Val Leu Phe Val Phe Ser Ser Ile

-20 -15 -10

WO 99/06550 PCT	Г/IB98/0
CCC CTT ACA TTT CTA TTT CAG AAA TTG CCT TTT GTT TGG ATT KGA GAA Pro Leu Thr Phe Leu Phe Gln Lys Leu Pro Phe Val Trp Ile Xaa Glu -5 1 5 10	96
GAG ACT TTG GAG ACA TGG TAT TTG AAG AGC TGG Glu Thr Leu Glu Thr Trp Tyr Leu Lys Ser Trp 15 20	129
(2) INFORMATION FOR SEQ ID NO: 124:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 352 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Cancerous prostate	
 (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 293346 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.8 seq LSIFSLVLPVCRM/HR 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 124:	
ACAATTCCAG CTTATGTGTC CCTTTTATAA ACTTGTGATA CATTTTAACT GTGTATACAC	60
ATCTCTTGCC TCTATTGGTA GAGAGTATCT GSCAKGCCTA GCATGTGCTG GATGTCATAT	120
CAGATACTCA GTGTTATTTA TTGGGCTTAC AGTGATAACC AAAGCTCACA TGTTTTAGCA	180
CTCCCACTTC CATAAAGTGG AAGATGTCCC CTCTGCCTCT TCTCTCATCC CTCCTCAAAG	240
CAGCAGGAGT GACTTACCTG ATTGACCAGT TTAAGACTAT ATCTGAGCAG GC ATG CCA Met Pro	298
CAG TAC TGT CTC AGC ATC TTC TCT CTT GTG CTG CCT GTC TGC AGG ATG Gln Tyr Cys Leu Ser Ile Phe Ser Leu Val Leu Pro Val Cys Arg Met -15 -5	346
CAC AGG His Arg	352
(2) INFORMATION FOR SEQ ID NO: 125:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 194 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE	

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 15..143

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 4.8

seq LLAFGTSCSVVLY/DP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 125:

GACCAGTTGG CGAC ATG GTG GCA CCC GTG CTG GAG ACT TCT CAC GTG TTT 50 Met Val Ala Pro Val Leu Glu Thr Ser His Val Phe -40

TGC TGC CCA AAC CGG GTG CGG GGA GTC CTG AAC TGG AGC TCT GGG CCC 98 Cys Cys Pro Asn Arg Val Arg Gly Val Leu Asn Trp Ser Ser Gly Pro -30

AGA GGA CTT CTG GCC TTT GGC ACG TCC TGC TCC GTG GTG CTC TAT GAC 146 Arg Gly Leu Leu Ala Phe Gly Thr Ser Cys Ser Val Val Leu Tyr Asp -15

· CCC CTG GGT TGT TGT TAC CAA CTT GAA TGG TCA CAC CGC CCG TTC CGG 194 Pro Leu Gly Cys Cys Tyr Gln Leu Glu Trp Ser His Arg Pro Phe Arg

(2) INFORMATION FOR SEO ID NO: 126:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 346 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 134..247
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8

seq LSWLITWFGHXLS/DF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 126:

TTATCTACCC ACCACCTCAG GGATTTTATG GATCCAVCAA TGGRACAACA CCAMGCATAT	120
ATTAAACTAT CTG ATG CCC ATC ATT GAC CAG GTG AAT CCA GAG CTC CAT Met Pro Ile Ile Asp Gln Val Asn Pro Glu Leu His -35 -30	169
GAC TTC ATG CAG AGT GCT GAG GTA GGG ACC ATC TTT GCC CTC AGC TGG Asp Phe Met Gln Ser Ala Glu Val Gly Thr Ile Phe Ala Leu Ser Trp -25 -20 -15	217
CTC ATC ACC TGG TTT GGG CAT GWM CTG TCT GAC TTC AGG CAC GTC GTG Leu Ile Thr Trp Phe Gly His Xaa Leu Ser Asp Phe Arg His Val Val -10 5	265
CGG TTA TAT GAC TTC TTC CTR GCC TGC CAC CCA CTG ATG CCG ATT TAC Arg Leu Tyr Asp Phe Phe Leu Ala Cys His Pro Leu Met Pro Ile Tyr 10 15 20	313
TTT GCA GCC GTG ATT GTG TTG TAT CGC GAG CAG Phe Ala Ala Val Ile Val Leu Tyr Arg Glu Gln 25 30	346
(2) INFORMATION FOR SEQ ID NO: 127: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 374 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Cancerous prostate (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 63209 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.7 seq GICVLVPCSXSXX/WR	
AAKTKKKKGG AGCATTTCCT TCCCTGACAG CCGGACCTGG KACTGGGCTG GGGCCCTGGC	60
GG ATG GAG ACA TKC TGC CCC TGC TGC TGC TGC CCC TGC KGT GGG GDN Met Glu Thr Xaa Cys Pro Cys Cys Cys Pro Cys Xaa Gly Xaa -45 -40 -35	107
GGG TCC CTG CAK GAK AAG CCA GTK TAC GAG CTG CAA GTG CAG AAG TCG Gly Ser Leu Xaa Xaa Lys Pro Val Tyr Glu Leu Gln Val Gln Lys Ser -30 -25 -20	155
GTG ACG GTG CAG GAG GGC CTG TGC GTC CTT GTG CCC TGC TCC TKC TCT /al Thr Val Gln Glu Gly Leu Cys Val Leu Val Pro Cys Ser Xaa Ser	203

	97		
-15	-10	-5	
TAS SCC TGG AGA TCC TGG T Xaa Xaa Trp Arg Ser Trp T 1	AT TCC TCT CCC CCA yr Ser Ser Pro Pro 5	CTC TAC GTC TAC TGG Leu Tyr Val Tyr Trp 10	251
TTC CGG GAC GGG GAG ATC Corphe Arg Asp Gly Glu Ile P 20	CA TAC TAC GCT GAG ro Tyr Tyr Ala Glu 25	GTT GTG GCC ACA AAC Val Val Ala Thr Asn 30	299
AAC CCA GAC AGA AGA KTG A Asn Pro Asp Arg Arg Xaa L 35			347
CCT TGG GGA TGT CCA GAA G Pro Trp Gly Cys Pro Glu G 50	· ·		374
(2) INFORMATION FOR SEQ I (i) SEQUENCE CHARACT (A) LENGTH: 39 (B) TYPE: NUCL (C) STRANDEDNE (D) TOPOLOGY: (ii) MOLECULE TYPE:	TERISTICS: 99 base pairs LEIC ACID ESS: DOUBLE LINEAR		
(vi) ORIGINAL SOURCE (A) ORGANISM:	E:	ate	
(ix) FEATURE: (A) NAME/KEY: (B) LOCATION: (C) IDENTIFICA (D) OTHER INFO	295345 ATION METHOD: Von E ORMATION: score 4.	_	
-(xi) SEQUENCE DESCR	IPTION: SEQ ID NO:	128:	
ATTTTCAGTG CAGCCTGCCA GAC	CTCTTCT GGAGGAAGAC	TGGACAAAGG GGGTCACACA	60
TTCCTTCCAT ACGGTTGAGC CTC	TACCTGC CTGGTGCTGG	TCACAGTTCA GCTTCTTCAT	120
GRWKGGTGGA TCCCAATGGC AAT	GAATCCA GTGCTACATA	CTTCATCCTA ATAGGCCTCC	180
CTGGTTTAGA AGAGGCTCAG TTC	TGGTTGG CCTTCCCATT	GTGCTCCCTC TACCTTATTG	240
CTGTGCTAGG TAACTTGACA ATC	ATCTACA TTGTGCGGAC	TGAGCACAGC CTGC ATG	297
AGC CCA TGT ATA TAT TTC T Ser Pro Cys Ile Tyr Phe P -15			345
TCT CCA CCT CAT CCA TGC C Ser Pro Pro His Pro Cys P			393

V	VO 99/06550		. 98	PC	T/IB98/01232
1		5	10	15	
	CTA Leu				399
(2)	INFORMATION	FOR SEQ ID NO:	129:		
	(A) (B) (C)	CNCE CHARACTERIST LENGTH: 110 base TYPE: NUCLEIC AC STRANDEDNESS: DO TOPOLOGY: LINEAR	e pairs CID DUBLE		
	(ii) MOLE	CULE TYPE: CDNA			
	(A)	INAL SOURCE: ORGANISM: Homo S TISSUE TYPE: Nor	Sapiens mal prostate		
	(B) (C) (D)	NAME/KEY: sig_pe LOCATION: 1292 IDENTIFICATION M OTHER INFORMATIO	! IETHOD: Von Heijne N: score 4.7 seq VLKCLSFSX		
	(xi) SEQUI	ENCE DESCRIPTION:	SEQ ID NO: 129:		
AAGC	AACCGG G AT(Me	G GGA CGG GGA GAG t Gly Arg Gly Glu -25	G AGG AGG CAC TAC Arg Arg His Tyr -20	TGG GGA CCT AAG Trp Gly Pro Lys -15	50
CTG (GTT CTC AAA Val Leu Lys	TGC CTC TCC TTT Cys Leu Ser Phe -10	TCS SCT CCA AGC C Ser Xaa Pro Ser I -5	CTC CCA GGC TTC Leu Pro Gly Phe	98
	FGG TCC CTA Frp Ser Leu 5				110
(2)	NFORMATION	FOR SEQ ID NO: 1	30:		
	(A) (B) (C)	ICE CHARACTERISTIC LENGTH: 251 base TYPE: NUCLEIC ACT STRANDEDNESS: DOU TOPOLOGY: LINEAR	pairs ID		
	(ii) MOLEC	CULE TYPE: CDNA			
	(A)	NAL SOURCE: ORGANISM: Homo Sa TISSUE TYPE: Cand	apiens cerous prostate		
-	(ix) FEATU (A)	RE: NAME/KEY: sig_per	otide		•

(B) LOCATION: 9..164

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 4.7

seg LLAKALHLLKSSC/AP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 130:

AGCCTGCG ATG TCT CAA GAT GGC GGA STG GGC GAA TTA AAG CAC ATG GTG

Met Ser Gln Asp Gly Gly Xaa Gly Glu Leu Lys His Met Val

-50

ATG AGT TTC CGG GTG TCT GAG CTC CAG GTG CTT CTT GGC TTT GCT GGC

Met Ser Phe Arg Val Ser Glu Leu Gln Val Leu Leu Gly Phe Ala Gly

-35 -30 -25

CGG AAC AAG AGT GGA CGG AAG CAC GAG CTC CTG GCC AAG GCT CTG CAC

Arg Asn Lys Ser Gly Arg Lys His Glu Leu Leu Ala Lys Ala Leu His

-20

-15

-10

CTC CTG AAG TCC AGC TGT GCC CCT AGT GTC CAG ATG AAG ATC AAA GAG
Leu Leu Lys Ser Ser Cys Ala Pro Ser Val Gln Met Lys Ile Lys Glu
-5 1 5 10

CTT TAC CGA CGA CGC TTT CCC CGG AAG ACC CTG GGG CCC TCT GAT CTC
Leu Tyr Arg Arg Arg Phe Pro Arg Lys Thr Leu Gly Pro Ser Asp Leu
15 20 25

TCC CTA AAG 251 Ser Leu Lys

(2) INFORMATION FOR SEQ ID NO: 131:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 272 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 18..224

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 4.6

seq LGPSLSSLPSALS/LM

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 131:

TATTTGGCCC CAAGCCG ATG CAT CAC AGG ATG AAT GAA ATG AAC CTG AGT 50

Met His His Arg Met Asn Glu Met Asn Leu Ser

-65

										-							
CCA Pro	GTG Val	GGG Gly	ATG Met -55	GAG Glu	CAG Gln	CTG Leu	ACT Thr	TCA Ser -50	TCC Ser	TCT Ser	GTG Val	AGC Ser	AAT Asn -45	GCC Ala	TTG Leu	98	
CCA Pro	GTC Val	TCA Ser -40	GGA Gly	AGT Ser	CAC His	CTG Leu	GGA Gly -35	TTG Leu	GCT Ala	GCC Ala	TCA Ser	CCC Pro -30	ACT Thr	CAC His	AGT Ser	146	
GCC Ala	ATC Ile -25	CCT Pro	GCC Ala	CCA Pro	GGC Gly	CTC Leu -20	CCA Pro	GTG Val	GCA Ala	ATT Ile	CCA Pro -15	AAC Asn	CTG Leu	GGT Gly	CCC Pro	194	
TCC Ser -10	CTG Leu	AGC Ser	TCT Ser	CTG Leu	CCT Pro -5	TCT Ser	GCT Ala	CTG Leu	TCT Ser	TTA Leu 1	ATG Met	CTA Leu	CCA Pro	ATG Met 5	GGT Gly	242	
DTT Xaa	GGG Gly	GAT Asp	CGA Arg 10	GGG Gly	GTG Val	ATG Met	TGT Cys	GGG G1 y 15	TTA Leu							272	

(2) INFORMATION FOR SEQ ID NO: 132:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 127 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 62..118
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.6

seq IWNLFSLFSTSTT/LP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 132:

ACATCCTTGA TTCTTTACTT TCTCTTAACA CCCTGTATCC AGCTGGTCAT AAATCTAGCA 60

G ATG CTA CAT TCA GAT AAC ATC TGG AAT CTA TTT TCC CTA TTT TCT ACT 109

Met Leu His Ser Asp Asn Ile Trp Asn Leu Phe Ser Leu Phe Ser Thr

-15

-10

TCT ACT ACC CTG CCC CGG Ser Thr Thr Leu Pro Arg

127

- (2) INFORMATION FOR SEQ ID NO: 133:
 - (i) SEQUENCE CHARACTERISTICS:

		(B) TYP (C) STR	GTH: 135 E: NUCLE ANDEDNES: OLOGY: L	IC ACID S: DOUBL								
	(ii) M	OLECULE	TYPE: C	DNA								
	(vi) O	(A) ORG	SOURCE: ANISM: HO SUE TYPE			state	:					
	 (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 475 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.6 seq FHSAAGWSGGGQA/CG 											
	(xi) S	EQUENCE	DESCRIP!	TION: SE	Q ID	NO:	133:					
ATT	ATG CAA Met Gln	CCC GCC Pro Ala	TCC CCG Ser Pro -20	CCC GCC Pro Ala	CGG Arg	TGG Trp -15	AGC Ser	TTC Phe	CAC His	TCG Ser	GCT Ala -10	48
GCG Ala	GGC TGG Gly Trp	AGC GGC Ser Gly -5	Gly Gly	CAG GCG Gln Ala	TGC Cys 1	GGA Gly	GGA Gly	CAC His	TCC Ser 5	TGC Cys	GAC Asp	96
CAG Gln	GTA CTG Val Leu 10	GCT GTG Ala Val	ATC GAA Ile Glu	CTT CTC Leu Leu 15	AAC Asn	CCT Pro	CTC Leu	AGG Arg 20				135
(2)	INFORMAT	ION FOR	SEQ ID	NO: 134:								
		(A) LEN (B) TYP (C) STR	CHARACTEI GTH: 233 E: NUCLEI ANDEDNESS OLOGY: LI	base pa IC ACID S: DOUBL								
	(ii) M	OLECULE	TYPE: CI	DNA								
		(A) ORG	SOURCE: ANISM: Ho SUE TYPE			rost	ate					
		(B) LOC (C) IDE	E/KEY: s: ATION: 1: NTIFICAT: ER INFORM	38191 ION METH	DD: V	on Hee 4.	5					
	(xi) S	EQUENCE	DESCRIP	TION: SE	Q ID	NO:	134:					

			102			
ACCTTTCTGC	CACAGATGAC	GGAAACATTT	AAAGTTATGG	ATTGTGTCTC	TGCATCCTCT	120

TCCCTTCACA CCAGCCA ATG TGT TTT TCA TTT CTC TTG GCT GGC TCA ATT

Met Cys Phe Ser Phe Leu Leu Ala Gly Ser Ile

-15

TCC CAC ATG TTC TCC CAA GCT CTT CCT CTC CAC TCC CCA GGG CTT CCC Ser His Met Phe Ser Gln Ala Leu Pro Leu His Ser Pro Gly Leu Pro

ACC ACA AAC CGC ACG
Thr Thr Asn Arg Thr
10

(2) INFORMATION FOR SEQ ID NO: 135:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 214 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 137..199
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.5

seq SILFHCSVCLFLC/QY

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 135:

ATATGGCAAG AGATAGAGAT CTAGTTTCAT TCTTCTGCAT ATGGATATCC AATTTTCCCA 60

GCACCATTTA TTGAAGAGAC AGTCCTTTTG CCAGTKTATG TTCTTGGCAA CTTTGTTGAA 120

AATGCATTTA CTGTAG ATG TAT GGA TTC ATT ATT GGG TTA TCT ATT CTG TTC 172

Met Tyr Gly Phe Ile Ile Gly Leu Ser Ile Leu Phe -15

CAT TGT TCT GTG TGT CTG TTT TTA TGC CAG TAC CAT GCC TGG
His Cys Ser Val Cys Leu Phe Leu Cys Gln Tyr His Ala Trp

-5

1

5

(2) INFORMATION FOR SEQ ID NO: 136:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 231 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR

50

(ii)	MOLECULE TYPE: CDNA	
(vi)	ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate	
(ix)	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 139210 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.5 seq SLLGCXLAININT/FP	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 136:	
ATCCTATTGT	GTCGTGTAGC TTGTTCTCTA TTTTATAGGT CATTTAAAAT AAAACTCACC	60
TTTGACTTTG	TTTAGTCTCT GTTACATGTT TGCTTTTTGT TTCGTTTATG TTTGTACATT	120
TCTCATGTKT	TTCTKKCT ATG TCT TTT GGT KGT ATT CTA ACT TTT AGA GTC Met Ser Phe Gly Xaa Ile Leu Thr Phe Arg Val -20 -15	171
TCT TTA TTO Ser Leu Le	G GGA TGT CNT CTA GCG ATA AAT ATA AAT ACA TTT CCC TCT u Gly Cys Xaa Leu Ala Ile Asn Ile Asn Thr Phe Pro Ser -10 -5 1	219
AAC AAC CA Asn Asn Hi		231
(2) INFORM	ATION FOR SEQ ID NO: 137:	
-	SEQUENCE CHARACTERISTICS: (A) LENGTH: 269 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
	MOLECULE TYPE: CDNA	
(V1)	ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Prostate	
(ix)	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 1277 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.4 seq LGRLCAGSSGVXG/AR	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 137:	

AAAAGCGAGC C ATG GCT GTC TAC GTC GGG ATG CTG CGC CTG GGG AGG CTG Met Ala Val Tyr Val Gly Met Leu Arg Leu Gly Arg Leu

WO 99/06550		1	04	P	CT/IB98/01232		
	-20		-15		-10)	
TGC GCC GGG AGC TCC Cys Ala Gly Ser Ser -5	Gry var Xaa	GGG GCC Gly Ala	CGG GCC Arg Ala	GSC CTC Xaa Leu 5	TCT CGG Ser Arg	98	
AGT TGG CAG GAA GCC Ser Trp Gln Glu Ala 10	AGG TTG CAG Arg Leu Gln 15	GGT GTC Gly Val	CGC TTC Arg Phe	CTC AGT Leu Ser 20	TCC AGA Ser Arg	146	
GAG GTG GAT CGC ATG Glu Val Asp Arg Met 25	GTC TCC ACG Val Ser Thr 30	CCC ATC Pro Ile	GGA GGC Gly Gly 35	CTC AGC Leu Ser	TAC GTT Tyr Val	194	
CAG GGG TGC ACC AAA Gln Gly Cys Thr Lys 40	AAG CAT CTT Lys His Leu 45	AAC AGC Asn Ser	AAG ACT Lys Thr	GTG GGC Val Gly	CAG TGC Gln Cys 55	242	
CTG GAG ACC ACA GCA Leu Glu Thr Thr Ala 60	CAG AGG GTC Gln Arg Val	CCG Pro				269	
(2) INFORMATION FOR SEQ ID NO: 138: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 276 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Hypertrophic prostate (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 187255 - (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.4 seq LVSIFFFWEVTNA/FL (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 138:							
AGATAATTTT GATGAAACC							
CCTAGTGTTT TWGTTTATK							
GACCACCAAA TAGTTTAAT						180	
	nr lle Tyr Le 20	eu Val II -1	e Ser Le	u Val Se	r Ile -10	228	
TTT TTC TTT TGG GAA Phe Phe Phe Phe Trp Giu	GTA ACT AAT (Val Thr Asn A	GCT TTC C la Phe I l	CTT AAG GO eu Lys A	CC AGG C la Arg A 5	GT TGG rg Trp	276	

(2)	INFORMAT	FION FOR SEQ ID NO: 139:	
	(i) SE	EQUENCE CHARACTERISTICS: (A) LENGTH: 137 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
	(ii) M	MOLECULE TYPE: CDNA	
	, ,	ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate	
		TEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 36101 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.4 seq SLPLTTGSSWSLS/SQ	
	(xi) S	SEQUENCE DESCRIPTION: SEQ ID NO: 139:	
ACC'	TTCTCAA G	GAACTGTGTT CACCCACTTC CCCAC ATG GCC CTT CCA CCC AAG Met Ala Leu Pro Pro Lys -20	53
GGA Gly	TGT GGT Cys Gly -15	AGT CTC CCT TTG ACT ACT GGG TCT TCC TGG AGC CTT TCT Ser Leu Pro Leu Thr Thr Gly Ser Ser Trp Ser Leu Ser -10	101
TCT Ser 1	CAA ATA Gln Ile	GGA AGC CCT GCT ATT TCC AAC CCT AGG Gly Ser Pro Ala Ile Ser Asn Pro Arg 5 10	137
(2)	INFORMAT	CION FOR SEQ ID NO: 140:	
		QUENCE CHARACTERISTICS: (A) LENGTH: 127 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
	(ii) M	OLECULE TYPE: CDNA	
	•	RIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Hypertrophic prostate	
	•	CEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 4491 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.3 seq FLSWASFLAPLLR/SP	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 140:

(MI, DESCRIPTION: SEQ ID NO: 140:	
GTCATTTGTC CGTTTCTTCC CCCTTGCCAA TTTTTTAATT AGA ATG TTT GTC TTT Met Phe Val Phe -15	55
TTG TCT TGG GCA AGT TTC TTA GCC CCT CTA CTG AGG AGC CCA TTT CTT Leu Ser Trp Ala Ser Phe Leu Ala Pro Leu Leu Arg Ser Pro Phe Leu -10 -5 1	103
CAT TGT CTA ATG GGG ATG CCA GGG His Cys Leu Met Gly Met Pro Gly 5 10	127
(2) INFORMATION FOR SEQ ID NO: 141: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 302 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 150233 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.3 seq LLSCSPLXPLGKS/GF (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 141:	
AAKAGTCAGC AGGAGTKAGT TCAGGAATCC TCGGGACAAG GCACTTTCCT GAGCACTGGA	
CCAGCGACCT CTTGGCTTCC AGGGAGGACA CACAGCCATC ATGGWACCCA THTCTCAGAA	60
GAGTCCAGGC AAACAGTTTA CATTTTCTT ATG AWA ATG AAG TCT GCA AAC AAG Met Xaa Met Lys Ser Ala Asn Lys -25	173
ATT ACT TTA TTA ART CAC CAC CTT CTC AGC TGT TCT CCT CTG TGW CCT Ile Thr Leu Leu Xaa His His Leu Leu Ser Cys Ser Pro Leu Xaa Pro -20 -15 -10 -5	221
CTT GGA AAA AGC GGT TTT TCA TCC TGT CAA AGG CTG GGG AAA AGA GCT Leu Gly Lys Ser Gly Phe Ser Ser Cys Gln Arg Leu Gly Lys Arg Ala 1 5 10	269
TTA GTC TTT CCT ATT ATR AAG NCC ATC ACC Leu Val Phe Pro Ile Xaa Lys Xaa Ile Ile Thr	302

(2) INFORMATION FOR SEQ ID NO: 142:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 251 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Cancerous prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 150245 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.2</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 142:	
AATTTGATAA CATCAGCTAA TATTTTTCAA AGTTAGATTT TTGAGGTATA ATTTACATAA	60
GAGTTACTCT TTCTAGAGGT ATAGTTGAAT GCATTTTCAC AAATGTGTAC AATTGGATAA 1	.20
CCACCAMCAT WAWTCTAGAW ATATAGGTA ATG TGT AAT TAT AAT ATA TAT GTA Met Cys Asn Tyr Asn Ile Tyr Val -30 -25	.73
CTA TAT AAT ATA GGA TAT TTA TAC CAC CCA AAA AGT TTT CTC TTG CTT Leu Tyr Asn Ile Gly Tyr Leu Tyr His Pro Lys Ser Phe Leu Leu -20 -15 -10	21
TTT ATA GTC ATT CCC CAA ACC CCA CGT CCG Phe Ile Val Ile Pro Gln Thr Pro Arg Pro -5 1	251
(2) INFORMATION FOR SEQ ID NO: 143:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 383 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR 	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide</pre>	

(B) LOCATION: 84..164

- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4.2 seq PLLAAPLLRSLLP/RX

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 143:

AACTGAACAG CGGASCGGAC GGGGATCGCC GGCGGGGCGC AAGCGGAGGC GGCCCAGRGC CCGGCGGTCT CCGAGATGTC ACG ATG GCT GTG GCC ATG GTC AAA CTG TGT GAA 113 Met Ala Val Ala Met Val Lys Leu Cys Glu AGA GCG GGT CTG CCG CTA CTT GCT GCA CCA CTA CTT AGG TCA CTT CTT Arg Ala Gly Leu Pro Leu Leu Ala Ala Pro Leu Leu Arg Ser Leu Leu 161 -15 -10 CCA AGA GMA CCT CAG CCT GGA CCA GCT CAG CCT CGA TCT GTA CAA GGG 209 Pro Arg Xaa Pro Gln Pro Gly Pro Ala Gln Pro Arg Ser Val Gln Gly CAG CGT TGC CCT GCG AGA CAT CCA CCT GGA AAT CTG GTC TGT GAA CGA 257 Gin Arg Cys Pro Ala Arg His Pro Pro Gly Asn Leu Val Cys Glu Arg GGT GCT RGA GTC AAT GGA GTC ACC GCT GGA GCT RGT GGA ARG CTT CGT 305 Gly Ala Xaa Val Asn Gly Val Thr Ala Gly Ala Xaa Gly Xaa Leu Arg 35 40 GGG CTC CAT CGA GGT DGC CGT GCC CTG GGC TGC TCT GCT CAC CGA CCA Gly Leu His Arg Gly Xaa Arg Ala Leu Gly Cys Ser Ala His Arg Pro

383

(2) INFORMATION FOR SEQ ID NO: 144:

MTG CAC AGT GCG CGT GTC CGG CCT CCA GCT

Xaa His Ser Ala Arg Val Arg Pro Pro Ala

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 479 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:

65

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 99..464
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4.2 seq DVLLGLLKDVLLA/RP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 144:

TAAACTTCTG AAAGAAAGAG	AAGATCTTCC TATATGO	GAAA GAAAAATACT CCTTTATGGA	60
GAACCTGCTT CAAAATCAAA	TCGTGATTGT TTCAGGA	AG ATG CTA AAT GTG GTA AGA 1 Met Leu Asn Val Val Arg -120	16
		TAT TGT CTT TCC ATC CAC 1 Tyr Cys Leu Ser Ile His -105	64
	l Ile Cys Thr Gln	GTC CAC AAG CAG ACT GTG 2 Val His Lys Gln Thr Val -90 -85	12
		ATG GAT GTT AAC ATT GGT Met Asp Val Asn Ile Gly -70	60
		AAC TGC TGT ACC AAC GAA Asn Cys Cys Thr Asn Glu -55	80
		CTG CAA AGA GAA ATG ATG Leu Gln Arg Glu Met Met -40	56
		ATC ATC TTA GAT GAT ATT Ile Ile Leu Asp Asp Ile -25	04
	a Thr Asp Val Leu	CTT GGA CTT CTT AAA GAT 4 Leu Gly Leu Leu Lys Asp -10 -5	52
GTT TTA CTA GCA AGA CC Val Leu Leu Ala Arg Pr 1		4	79

(2) INFORMATION FOR SEQ ID NO: 145:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 208 base pairs
 - (3) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 107..187
 - (C) IDENTIFICATION METHOD: Von Heijne matrix

110	
(D) OTHER INFORMATION: score 4.2 seq AGLCIGSTSYVHG/DI	
(xi) SEQUENCÉ DESCRIPTION: SEQ ID NO: 145:	
ATTGGGAGCA GCAGCATCTA CTTCACAGAC CAGTGTCCAG TTAATTGTGT TTGTGGCAAT	60
CATCCTACAT AAGGCACCAG CTGCTTTTGG ACTGGTTTCC TTCTTG ATG CAT GCT Met His Ala -25	115
GGC TTA GAG CGG RAW TCG AWT CAG AAA GCA CTT GCT GGT CTT TGC ATT Gly Leu Glu Arg Xaa Ser Xaa Gln Lys Ala Leu Ala Gly Leu Cys Ile -20 -15 -10	163
GGC AGC ACC AGT TAT GTC CAT GGT GAC ATA CTT AGG ACT GAG CGG Gly Ser Thr Ser Tyr Val His Gly Asp Ile Leu Arg Thr Glu Arg	208
(2) INFORMATION FOR SEQ ID NO: 146:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 285 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Prostate</pre>	
<pre>(ix) FEATURE:</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 146:	
AATTGCTGGG CTCGAAGCAC AGGAGAGACC AGTCCTTCCT TGTCTCCACT GGGCTGTKTA	60
GTGCTTCTTT CCCAAGGACK TCCATCCCTT CCCCAGGCTT TATGGTTCCA GTKCTTCTAC	120
CATTCTGGAA GCTCCCTAGA ATCTCCTGGA ATG CTT AAT GGA CCT TTC CAG CAC Met Leu Asn Gly Pro Phe Gln His -35	174
CGA AAT TCA AGA ATT ATG ACT CAT CGG TCA GCA GAA AAG ACC CTG CTG Arg Asn Ser Arg Ile Met Thr His Arg Ser Ala Glu Lys Thr Leu Leu -25 -20 -15	222
GGA TCT TTG AGC TTG TGG AGG TGG TCG GCA ATG GAA CCT ACG GAC AGG Gly Ser Leu Ser Leu Trp Arg Trp Ser Ala Met Glu Pro Thr Asp Arg -10 -5 1 5	270

WO 99/06550 PCT/IB98/01232

	ACA .													285
(2)	INFO	RMAI	CION	FOR	SEQ	ID N	NO: 1	L47:						
	(i) SE	(A) (B) (C)	ICE C LENG TYPE STRA TOPO	TH: : NU NDEC	409 CLEI NESS	base C AC : DC	pai ID UBLE						
	(i.	i) M	OLEC	ULE	TYPE	: CE	NA							
	(v.	i) C	(A)	NAL ORGA TISS	NISM	: Ho		_		rost	ate			
	(i:	x) F	(B) (C)	NAME LOCA	TION TIFI	: 44 CATI	17 ON M	5 ETHC	D: V	e 4.	leijn l SLTCQ			
	(x:	i) S	EQUE	NCE	DESC	RIPT	CION:	SEC) ID	NO:	147:			
AAGO	STTGT	AG #	CGC1	GCGC	SC CC	CGGCC	CCGGC	GG(TAA	AATA	CAG		GTG Val	55
	CCA Pro													103
	GTA Val													151
	TGC (199
	ATA Ile 10													247
	AGA Arg													295
	CTC Leu													343
	TCA Ser													391

ATA	ATT	AAT	CTG	AGC	ACA
Ile	Ile	Asn	Leu	Ser	Thr
		75			

409

(2)	INFORMATION	FOR	SEQ	ID	NO:	148:
(2)	INFORMATION	FOR	SEQ	ΙD	NO:	148:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 279 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 184..267
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4

seq FSLLALSMLKGTG/KV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 148:

ACATAATCGG CCTTTATGTT ACACTGCCTG GCCAGCCCCT GTTATTCTAG TGCATAATTG 60

ATGGTGCTCA CAAGTGGAAA AGTTAGAAAA GCGGAAGTAA TGTGACGCAG CAGTGCCATG 120

RAGCSSCCGG DVCCCCGGCA GTGAGGGCAA TGCAGAGATG GGCTGCTGCT GGCTACCGCC 180

AGG ATG CCT CAG AAG GGC CTG GGC TTA CTT GGC ATC TTG TCA GGA GAC 228

Met Pro Gln Lys Gly Leu Gly Leu Gly 1le Leu Ser Gly Asp -15

TTT TCC CTT CTT GCT TTG TCC ATG CTG AAA GGG ACA GGA AAG GTA GGC 276

Phe Ser Leu Leu Ala Leu Ser Met Leu Lys Gly Thr Gly Lys Val Gly -5

GGG Gly

ACAGGA AAG GTA GGC 279

GGGG Gly

(2) INFORMATION FOR SEQ ID NO: 149:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 326 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 69..233
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4

seq AALCGISLSQLFP/EP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 149:

AAGAACCTGA GCAGCCTGTC TTCAGACAGA GAGAGGCCCA CGGCTGTTTC TTGAAAYTGG 60 CGCTGGGA ATG GCC ATG TGG AAC AGG CCA TGB BAG ANG CTG CCT CAG CAG 110 Met Ala Met Trp Asn Arg Pro Xaa Xaa Xaa Leu Pro Gln Gln -55 CCT CTS STA GCT GAG CCC ACT GCA GAG GGG GAG CCA CAC CTG CCC ACG 158 Pro Leu Xaa Ala Glu Pro Thr Ala Glu Gly Glu Pro His Leu Pro Thr -40 -35 GGC CGG GAS BYG ACT GAG GCC AAC CGC TTC GCC TAT GCT GCC CTC TGT 206 Gly Arg Xaa Xaa Thr Glu Ala Asn Arg Phe Ala Tyr Ala Ala Leu Cys -25 -20 -15GGC ATC TCC CTG TCC CAG TTA TTT CCT GAA CCC GAA CAC AGC TCC TTC 254 Gly Ile Ser Leu Ser Gln Leu Phe Pro Glu Pro Glu His Ser Ser Phe TGC ACA GAG TTC ATG GCA GGC CTG GTG SKM TGG CTG GAG TTG TCT GAA 302 Cys Thr Glu Phe Met Ala Gly Leu Val Xaa Trp Leu Glu Leu Ser Glu GCT GTC TTG CCA ACC ATG ACT GCT 326

(2) INFORMATION FOR SEO ID NO: 150:

Ala Val Leu Pro Thr Met Thr Ala

25

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 194 base pairs

30

- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 126..182
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4

seq LLLSPWVTVPVWS/SS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 150:

CCTAGTGCTT AAGGGGATTT AGCATCATCC AAGCAGGGTA AACTTTTGTT TTGTTAAAAG														
AAAAATGTGT TATTCAAGTT GGTGTCCCCA GTTGTAGCTA ACACATCTGG AATGCACTAA	120													
CCAAA ATG CTG TGC TTT GGA GAC CTG CTT TTG TCA CCG TGG GTA ACC GTT Met Leu Cys Phe Gly Asp Leu Leu Leu Ser Pro Trp Val Thr Val -15 -10 -5	170													
CCC GTC TGG TCC AGT AGC CCG TGG Pro Val Trp Ser Ser Pro Trp 1														
(2) INFORMATION FOR SEQ ID NO: 151:														
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 170 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR 														
(ii) MOLECULE TYPE: CDNA														
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate														
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 27107 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4</pre>														
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 151:														
AAGTTAGGTT TAAAGTTTCC TCATTA ATG CAG GAA AAT GCT CAT AAC CTG AGG Met Gln Glu Asn Ala His Asn Leu Arg -25 -20	53													
CTT TTC AAG TGT TTA TTA ATT TAC TTT CTG GGG CTG GCT GCT GAT ACT Leu Phe Lys Cys Leu Leu Ile Tyr Phe Leu Gly Leu Ala Ala Asp Thr -15 -10 -5	101													
TAT TTC AGA TCA AAG AGA AAG CCT GTG TCT TTC GTA GTT ACT GTG KKG Tyr Phe Arg Ser Lys Arg Lys Pro Val Ser Phe Val Val Thr Val Xaa 1 5 10	149													
CMA GGA AMC TAT GCC ACA GGG Xaa Gly Xaa Tyr Ala Thr Gly 15 20	170													

(2) IMFORMATION FOR SEQ ID NO: 152:

(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 315 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: DOUBLE(D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 127303 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 152:	
ACCAAGTCCT CCCAAGTTAT TAACTGGTCA AAAAGGMTTA AAGGMTTAGT TCTTAATAGT	60
PAAGATGCCA CCCATTCAGG GTTTTTTGCT TTCTAAGAGG GAACTTTTAC AGGCATAATT	120
GAGAGA ATG CAT ACA TGC TCT CTA CCT TGT CTT CTC TTT GCT CAG CTG Met His Thr Cys Ser Leu Pro Cys Leu Leu Phe Ala Gln Leu -55 -50	168
CTA GAA TTT TGT AGC TTT CCT CCA GAT GTG CCT CAT AAC TGT GCG CCT Leu Glu Phe Cys Ser Phe Pro Pro Asp Val Pro His Asn Cys Ala Pro -45 -35 -30	216
ATT GTC TCA GTC AGG CCG CCT AAT ATT GTA GCA GCC TTT GAA GGG TGC Ile Val Ser Val Arg Pro Pro Asn Ile Val Ala Ala Phe Glu Gly Cys -25 -20 -15	264
FCT GTA GCC ACT GCT CTT TTT CCT CCC TTG TGC ATC TCC ACA GGG AAT Ser Val Ala Thr Ala Leu Phe Pro Pro Leu Cys Ile Ser Thr Gly Asn -10 -5 1	312
GAG Glu	315
(2) INFORMATION FOR SEQ ID NO: 153:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 342 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR 	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:	

(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Hypertrophic prostate

(i	ix)	FEATURE:	
١, ٦		I DUI OUG:	

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 55..138
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4

seq PLLGVLFFQGVYI/VF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 153:

AGT	CGTT	ACC	GGGA	GCTG	TA A	ACAA	GGTG	T GC	AAGC	ATCT	GAA	.GAGC	TGC	CGGG	ATG Met	57
CAG Gln	CAG Gln	AGA Arg -25	GGA Gly	GCA Ala	GCT Ala	GGA Gly	AGC Ser -20	CG T Arg	GGC Gly	TGC Cys	GCT Ala	CTC Leu -15	TTC Phe	CCT Pro	CTG Leu	105
CTG Leu	GGC Gly -10	GTC Val	CTG Leu	TTC Phe	TTC Phe	CAG Gln -5	GGT Gly	GTT Val	TAT Tyr	ATC Ile	GTC Val 1	TTT Phe	TCC Ser	TTG Leu	GAG Glu 5	153
ATT Ile	CGT Arg	GCA Ala	GAT Asp	GCC Ala 10	CAT His	GTC Val	CGA Arg	GGT Gly	TAT Tyr 15	GTT Val	GGA Gly	GAA Glu	AAG Lys	ATC Ile 20	AAG Lys	201
TTG Leu	AAA Lys	TGC Cys	ACT Thr 25	TTC Phe	AAG Lys	TCA Ser	ACT Thr	TCA Ser 30	GAT Asp	GTC Val	ACT Thr	GAC Asp	AAG Lys 35	CTT Leu	ACT Thr	249
ATA Ile	GAC Asp	TGG Trp 40	ACA Thr	TAT Tyr	CGC Arg	CCT Pro	CCC Pro 45	AGC Ser	AGC Ser	AGC Ser	CAC His	ACA Thr 50	GTA Val	TCA Ser	ATA Ile	297
KTK Xaa	CAT His 55	TAT Tyr	CAG Gln	TCT Ser	TTC Phe	CAG Gln 60	TAC Tyr	CCA Pro	ACC Thr	ACA Thr	GCA Ala 65	GGC Gly	ACA Thr	TTT Phe		342

(2) INFORMATION FOR SEQ ID NO: 154:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 429 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 109..225
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.9

seq LILNRSLPTASSS/SS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 154:

AAA	ATGI	CAC :	rgaat	GTC	CA CT	TTGO	GCCI	A GGC	CTGGC	GCAC	CGA	GGAC	ACA (GGGG	AACTAA	. 60
GAC	ACAG1	rcc r	rggto	CACTO	G G	AAACT	CAC	A GCC	CTGTT	rggg	AAA	GAAA(1 GAV a Xaa	117
				TCA Ser												165
				CTC Leu												213
				TCC Ser 1												261
				ATG Met												309
				CCT Pro												357
				AGM Xaa												405
				GGT Gly 65												429

(2) INFORMATION FOR SEQ ID NO: 155:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 351 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (3) LOCATION: 1..350
 - (C) IDENTIFICATION METHOD: fasta
 - (D) OTHER INFORMATION: identity 99.1 region 18..366 id D83597

vrt

(i:	x)	FEATURE:
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- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 127..186
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 3.9 seq FFWVVLFSAGCKV/IT
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 155:

ATTTCTTGTT CCAAGATCAC CCTTCTGAGT ACCTCTCTGG CTGCCAAATT GCCAGGGCCT TCACAGTTTG ATTCCATTTC TCAGCTCCAA GCATTAGGTA AACCCACCAA GCAATCCTAG CCTGTG ATG GCG TTT GAC GTC AGC TGC TTC TTT TGG GTG GTG CTG TTT 168 Met Ala Phe Asp Val Ser Cys Phe Phe Trp Val Val Leu Phe -15 TCT GCC GGC TGT AAA GTC ATC ACC TCC TGG GAT CAG ATG TAC ATT GAG 216 Ser Ala Gly Cys Lys Val Ile Thr Ser Trp Asp Gln Met Tyr Ile Glu AAA GAA GCC AAC AAA ACA TAT AAC TGT GAA AAT TTA GGT CTC AGT GAA 264 Lys Glu Ala Asn Lys Thr Tyr Asn Cys Glu Asn Leu Gly Leu Ser Glu 15 20 ATC CCT GAC ACT CTA CCA AAC ACA ACA GAA TTT TTG GAA TTC AGC TTT Ile Pro Asp Thr Leu Pro Asn Thr Thr Glu Phe Leu Glu Phe Ser Phe 312 AAT TTT TTG CCT ACA ATT CAC AAT AGA ACC TCC AGC AGG 351 Asn Phe Leu Pro Thr Ile His Asn Arg Thr Ser Ser Arg 50

(2) INFORMATION FOR SEQ ID NO: 156:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 410 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 96..383
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.9
 - seq IMNLTVMLDTAXG/KX
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 156:

.

CTTTATTAAT TCTCACGCTG CGGCCCTGGA AAGCG ATG GAG GTG GCG GCT AAT 113 Met Glu Val Ala Ala Asn -95 · TGC TCC CTA CGG GTG AAG AGA CCT CTG TTG GAT CCC CGC TTC GAG GGT Cys Ser Leu Arg Val Lys Arg Pro Leu Leu Asp Pro Arg Phe Glu Gly TAC AAG BTC TCT CTT GAG CCG CTG CCT TGT TAC CAG CTG GAG CTT GAC 209 Tyr Lys Xaa Ser Leu Glu Pro Leu Pro Cys Tyr Gln Leu Glu Leu Asp -70 GCA GCT GTG GCA KAG GTA AAA CTT CGA GAT GAT CAA TAT ACA CTG GAA 257 Ala Ala Val Ala Xaa Val Lys Leu Arg Asp Asp Gln Tyr Thr Leu Glu -55 CAC ATG CAT GCT TTT GGA ATG TAT AAT TAC CTG CAC TGT GAT TCA TGG 305 His Met His Ala Phe Gly Met Tyr Asn Tyr Leu His Cys Asp Ser Trp -40 -35TAT CAA GAC AGT GTC TAC TAT ATT GAT ACC CTT GGA AGA ATT ATG AAT Tyr Gln Asp Ser Val Tyr Tyr Ile Asp Thr Leu Gly Arg Ile Met Asn TTA ACA GTA ATG CTG GAC ACT GCC TTW GGR AAA MCA CGA GAG GTG TTT 401 Leu Thr Val Met Leu Asp Thr Ala Xaa Gly Lys Xaa Arg Glu Val Phe -10

(2) INFORMATION FOR SEQ ID NO: 157:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 347 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:

CGA CTC CTA

Arg Leu Leu

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 63..179
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 3.9

seq VLAIGLLHIVLLS/IP

410

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 157:

AGGGAACCGA TCCCGGGCCG TTGATCTTCG GCCCCACACG AACAGCAGAG AGGGGCATCA 60
GG ATG AAT GTK GGC ACA GCG CAC AGS DAG GTG AAC CCC AAC ACG CGG 107

										20							
	Met	Asn	Val	Gly	Thr -35	Ala	His	Xaa		Val -30	Asn	Pro	Asn	Thr	Arg -25		
GTH Val	(ATC	AAC Asn	AGC Ser	CGT Arg -20	GTA	ATC Ile	TGG Trp	CTC Leu	TCC Ser -15	TAC Tyr	GTG Val	CTG Leu	GCC Ala	ATC Ile -10	GGT Gly	155	
CTC	CTC Leu	CAC His	ATC Ile -5	Val	CTG Leu	CTG Leu	AGC Ser	ATC Ile 1	CCG Pro	TTT Phe	GTK Val	AGT Ser 5	GTC V al	CCT Pro	GTC Val	203	
GTC Val	TGG Trp	ACC Thr	CTC Leu	ACC Thr	AAC Asn	CTC Leu 15	ATT Ile	CAC His	AAC Asn	ATG Met	GGC Gly 20	ATG Met	TAT Tyr	ATC Ile	TTC Phe	251	
CTG Leu 25	HIS	ACG Thr	GTG Val	AAG Lys	GGG Gly 30	WCA Xaa	CCC Pro	TTT Phe	GAG Glu	ACC Thr 35	CCG Pro	GAC Asp	CAG Gln	GGC Gly	AAG Lys 40	299	
GCG Ala	AGG Arg	CTG Leu	CTW Leu	WCC Xaa 45	CAC His	TGK Xaa	TDA Xaa	GCA Ala	GAT Asp 50	GGA Gly	TTA Leu	TGG Trp	GGT Gly	CCA Pro 55	GTT Val	347	
	(: (: (:i	ii) SE ii) M vi) C ix) F	(A) (B) (C) (D) (OLEC (A) (F) (EATU (A) (B) (C) (D)	TYPE STRATOPO CULE NAL ORGATISS NAME LOCA IDEN OTHE	TH: C: NU NDED LOGY TYPE SOUR NISM UE T /KEY TION TIFICR IN	151 CLEI NESS : LI :: CD CE: : Ho YPE: : Si : 8. CATI	base C AC : DO NEAR NA MO S Hyp G_pe .76 ON MI	pai ID UBLE apie ertr ptid ETHO	ns ophi e D: Ve score seq :	on H e 3. SWWT	eijn 9 LLSS	e ma					
\TTI	TTAT	ATG Met	GAA Glu	AAC Asn	TTT Phe -20	AAC . Asn i	ATG Met	TAT . Tyr	Lys .	AAT Asn -15	AAG Lys	AGC Ser	TGG Trp	TGG Trp	ACC Thr -10	49	
CTT Leu	TTG Leu	TCC Ser	TCA Ser	TCA Ser -5	CCC . Pro	AGC Ser	TTT . Phe i	ATG . Met	ATC . Ile :	AGT Ser	TTT Phe	GTT Val	TCA Ser 5	TCT Ser	GTA Val	97	
ATC Leu	CCA Pro	GTG Val 10	CTA Leu	CTT Leu	ACC . Thr	ATC :	TCT A Ser A	AGG ' Arg	TTC . Phe	ATT Ile	TTG Leu	AAG (Lys (CAA . Gln	ATC Ile	CCA Pro	145	

	wo	99/06	550						1	21					P	СТ/ІВ9
	C CAC C Glr 25	ו														15
(2)	INE	FORMA	ATION	FOF	SEÇ) ID	NO:	159:								
	(i) S	(B) (C)	LEN TYP STR TOP	GTH: E: N ANDE	351 UCLE DNES	bas IC A S: D	e pa CID OUBL	irs							
	(ii)	MOLE	CULE	TYP	E: C	DNA									
	(vi)	ORIG (A) (F)	INAL ORG TIS	ANIS	м: н	omo : No	Sapi rmal	ens pro	stat	e					
	(ix)	(B) (C)	URE: NAM LOC. IDEI OTHI	ATIO NTIF	N: 1 ICAT	42 ION 1	258 METH	OD: '	Von 1 re 3 VLA	. 9					
	(xi)	SEQUI	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	159	:				
AGA'	TTCG	GCC	GGAG	CTGC	CA G	CGGG	GAGG	C TG	CAGC	CGCG	GGT'	TGTT.	ACA (GCTG	CTGGAG	60
CAG	CAGC	GGC	CCCC	GCTC	CC G	GGAA	CCGK'	T CC	CGGG	CCGT	TGR	TCTT	CGG (ccc	ACACGA	120
ACA	GCAG	AGA I	GGGG	CAGC	AG G	ATG Met	AAT Asn	GTG Val	GGS Gly	ACA Thr -35	GND Xaa	CAC His	AGC Ser	GAG Glu	GTG Val -30	17]
AAC Asn	CCC Pro	AAC Asn	ACG Thr	CGG Arg -25	GTG Val	Met	Asn	Ser	Arg	GGG Gly	Ile	Trp	Leu	Ser	Tyr	219
GTG Val	CTG Leu	GCC Ala	ATC Ile -10	GGT Gly	CTC Leu	CTC Leu	CAC His	ATC Ile -5	GTG Val	CTC Leu	CTG Leu	AGC Ser	ATC Ile 1	CCG Pro	T TT Phe	267
GTG Val	AGT Ser 5	GTC Val	CCT Pro	GTC Val	GTC Val	TGG Trp 10	ACC Thr	CTC Leu	ACC Thr	AAC Asn	CTC Leu 15	ATT Ile	CAC His	AAC Asn	ATG Met	315
GGC Gly	ATG Met	TAT Tyr	ATC Ile	TTC Phe	CTG Leu	TAC Tyr	ACG Thr	GTG Val	AAG Lys	GGG Gly	ACA Thr					351

(2) INFORMATION FOR SEQ ID NO: 160:

20

(i) SEQUENCE CHARACTERISTICS:

25

- (A) LENGTH: 234 base pairs
- (B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 88129 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.8</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 160:	
AABGCTTCGT AGTGGAGGAA CGGGTTTGGC GTGTGGGACG CAGCTGCCTC TGTACTGGGG	60
AGTCACGGAG TCCCGGGCTC CAGGGAC ATG GCG GCG GCC TCT GCG GTG TCG GTG Met Ala Ala Ala Ser Ala Val Ser Val -10	114
CTG CTG GTG GCG GCG GAG AGG AAC CGG TGG CAT CGT CTC CCG AGC CTG Leu Leu Val Ala Ala Glu Arg Asn Arg Trp His Arg Leu Pro Ser Leu -5 1 5 10	162
CTC CTG CCG CCG AGG ACA TGG GTG TGG AGG CAA AGA ACC ATG AAG TAC Leu Leu Pro Pro Arg Thr Trp Val Trp Arg Gln Arg Thr Met Lys Tyr 15 20 25	210
ACA ACA GCC ACA GGA AGA AAC ATG Thr Thr Ala Thr Gly Arg Asn Met 30 35	234
(2) INFORMATION FOR SEQ ID NO: 161:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 46l base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Cancerous prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 177308 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.8 seq SGSGLSWARLSOS/RS</pre>	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 161:

ACTO	CTTTC	SCC .	ACCC:	rcag <i>i</i>	AG GO	CGAG	CTGT	G GA	AGCC'	ГТGA	CTC	rtag(GGC (CGTT	TAGA	A 60
CCG	GGCC	CTC	GGAC	CGGC	GG GG	GTTT(CTGC	A CG	rgga	ACCG	GAA	CATC	rga (GATG	ATCGSI	4 120
RGG	CCT	GTG (GAGT	GTGG	GG A	GCGC	GGA	TT(CTTT	CTTC	CCT	CGAG	GCC (CGTG	CC ATO	
														GTC Val		227
														AGT Ser		275
														ATC Ile		323
														GAG Glu 20		371
														CTC Leu		419
			ATC Ile													461

(2) INFORMATION FOR SEQ ID NO: 162:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 459 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 175..285
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.8
 - seq RPVLLHLHQTAHA/DE
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 162:

									12	4						
CAG	CCGA	GAC	TCAC	GGTC	AA G	CTAA	GGCG	A AG	AGTG	GGTG	GCT	GAAG	CCA '	TACT	ATTTTA	120
TAG.	AATT	AAT	GGRA	ARCMI	HG A	A AAGI	MCAT(C AC	AAAC	CAAG	AAG	AACT	TTG (GAAA	ATG Met	177
AAG Lys	CCT Pro -35	AGG Arg	AGA Arg	AAT Asn	TTA Leu	GAA Glu -30	GAA Glu	GAC Asp	GAT Asp	TAT Tyr	TTG Leu -25	CAT His	AAG Lys	GAC Asp	ACG Thr	225
GGA Gly -20	GAG Glu	ACC Thr	AGC Ser	ATG Met	CTA Leu -15	AAA Lys	AGA Arg	CCT Pro	GTG Val	CTT Leu -10	TTG Leu	CAT His	TTG Leu	CAC His	CAA Gln -5	273
ACA Thr	GCC Ala	CAT His	GCT Ala	GAT Asp 1	GAA Glu	TTT Phe	GAC Asp	TGC Cys 5	CCT Pro	TCA Ser	GAA Glu	CTT Leu	CAG Gln 10	CAC His	ACA Thr	321
CAG Gln	GAA Glu	CTC Leu 15	TTT Phe	CCA Pro	CAG Gln	TGG Trp	CAC His 20	TTG Leu	CCA Pro	ATT Ile	AAA Lys	ATA Ile 25	GCT Ala	GCT Ala	ATT Ile	369
ATA Ile	GCA Ala 30	TCT Ser	CTG Leu	ACT Thr	TTT Phe	CTT Leu 35	TAC Tyr	ACT Thr	CTT Leu	CTG Leu	AGG Arg 40	GAA Glu	GTA Val	ATT Ile	CAC His	417
CCT Pro 45	TTA Leu	GCA Ala	ACT Thr	TCC Ser	CAT His 50	CAA Gln	CAA Gln	TAT Tyr	TTT Phe	TAT Tyr 55	AAA Lys	ATT Ile	CAA Gln			459

(2) INFORMATION FOR SEQ ID NO: 163:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 141 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 25..81
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.7

seq IPCAHMLVCPTIG/DI

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 163:

AATTTGTAAG AATATTATAT ATAG ATG ATC ATC TGT TAT GAT ATT CCT TGT

Met Ile Ile Cys Tyr Asp Ile Pro Cys

-15

GCA CAT ATG TTG GTT TGT CCT ACT ATT GGT GAT ATT AAG TTT GAT CAC
Ala His Met Leu Val Cys Pro Thr Ile Gly Asp Ile Lys Phe Asp His

	WO 9	9/065	50				125 P								PCT	/IB98/01232
-10		*			-5					1				5		
TTG Leu	ATG Met	AAG Lys	TGG Trp 10	TAT Tyr	CCA Pro	TCA Ser	GAT Asp	TTC Phe 15	TCT Ser	ACT Thr	GAA Glu	AGG Arg	CTG Leu 20			141
(2)	INFO	ORMA'	поп	FOR	SEQ	ID N	10: 3	164:								
	()	.) SE	(A) (B) (C)	ICE C LENG TYPE STRA TOPC	TH: : NU NDEC	393 CLEI NESS	base C AC	e pai CID DUBLE								
	(i	.i) N	OLEC	CULE	TYPE	: CE	NA									
	7)	vi) C	(A)	NAL ORGA TISS	NISM	: Ho				rost	ate					
	(i	.x) F	(B) (C)	NAME LOCA IDEN	TION TIFI	: 18	42 ON M	40 ETHC	D: V	on H e 3. STLA	7					
	(>	i) S	EQUE	NCE	DESC	RIPT	'ION:	SEC) ID	NO:	164:					
AAC	CAGGO	CTC T	TTTA	AGAG	c co	GGTA	AGGG(G AGO	CGCAC	GNC	CAGA	ATACO	CTC F	AGCGC	TACCT	60
GGC	GAAC	TG (SATTI	CTC1	c co	GCC1	GCC	G GCC	CTGCC	CTGC	CACA	AGCCC	GA (тссе	CCACT	120
CCGC	TAGO	cc c	CATGO	CTG	SM AA	CCTC	TGAC	S ATT	RAGCA	ATA	тттт	TAGO	CAA C	TACT	TCAGT	180
GCG										GCC Ala -10						228
GCC Ala	ACC Thr	TTT Phe	GGG Gly	GCC Ala 1	GAT Asp	GAC Asp	TTG Leu	GTA Val 5	CTG Leu	ACC Thr	CTG Leu	AGC Ser	AAC Asn 10	CCC Pro	CAG Gln	276
										TGG Trp						324
										TGG Trp						372

393

GAG AAG AAG TAC GAC GCG

Glu Lys Asn Lys Tyr Asp Ala 45 50

	(i) S	(B) (C)	LEN TYP STR	CHAR GTH: E: N ANDE OLOG	263 UCLE DNES	bas IC A S: D	e pa CID OUBL								
	(ii)	MOLE	CULE	TYP	E: C	DNA									
	(vi)	ORIG (A) (F)	ORG	SOU! ANIS! SUE	м : н	omo :	Sapi rmal	ens pro	state	e					
	(.	ix)	(B) (C)	NAMI LOCA I DEI	E/KEY ATION NTIFI ER IN	N: 54	12 ION I	48 METHO	DD: '	Von E re 3 QLEO	. 7					
	(:	xi)	SEQUI	ENCE	DESC	CRIP'	rion	: SE	Q ID	NO:	165	:				
ACC	CTGA	ATA (CGAA	GAAC	AT A	AGCAZ	AAGC'	T AC	rgga	GACA	CCG	AGAA	CTA /	i	ATG Met -65	56
GGG Gly	GAA Glu	GAC Asp	CCT Pro	KCC Xaa -60	CAG Gln	CCC Pro	CGC Arg	AAG Lys	TAT Tyr -55	AAG Lys	AAG Lys	WWG Xaa	AAG Lys	AWG Xaa -50	GAG Glu	104
CTA Leu	CAG Gln	GGT Gly	GAT Asp -45	KGG Xaa	CCT Pro	CCC Pro	AGT Ser	TCT Ser -40	CCC Pro	ACT Thr	AAT Asn	GAT Asp	CCT Pro -35	ACC Thr	GTG Val	152
AAA Lys	TAT Tyr	GAG Glu -30	ACT Thr	CAG Gln	CCA Pro	CGG Arg	TTT Phe -25	ATC Ile	ACA Thr	GCC Ala	ACT Thr	GGA Gly -20	GGC Gly	ACC Thr	CTG Leu	200
CAC His	ATG Met -15	TAT Tyr	CAG Gln	TTG Leu	GAA Glu	GGG Gly -10	CTG Leu	AAC Asn	TGG Trp	CTA Leu	CGC Arg -5	TTC Phe	TCC Ser	TGG Trp	GCC Ala	248
			KWC Xaa													263
(2)	INFO	ORMA!	rion	FOR	SEQ	ID N	10: 3	166:								
	(i	l) SE	(B) (C)	LENG TYPE STRA	CHARA TH: NU NDED	372 CLEI NESS	base C AC : DC	pai CID OUBLE								
	(i	.i) N	40LEC	ULE	TYPE	: CD	NA			٠						
	(♥	vi) (ORIGI (A)		SOUR NISM		mo S	Sapie	ns							

151	TECHE	TVDE.	Prostate

(ix	.)	FEA	TURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 148..273
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 3.7

seq LLGCLQCCWLQSG/RA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 166:

ACCAATTTTG TAGTTATCTG ATCTGAAGGA AGATGTGTGT GGAGGTGTTT AGTGATGTTT 60 TCCGATGACG GTGATTCCCC CTAAATCTAC GTATTAAATA CAATGGAACA GGATCCACAG TTCACCCCTA ATAATATAGT TTACTGA ATG TTT TAT GTA GCT ATG ACC AAA ACT Met Phe Tyr Val Ala Met Thr Lys Thr -40 CAC AAA AGG ATC AGA AGC CTC TGT AAC ATC CAC CAT GGT TTG TTC CAG 222 His Lys Arg Ile Arg Ser Leu Cys Asn Ile His His Gly Leu Phe Gln -30 -25 TTT ACT CAG CAG CTC CTG GGC TGT CTT CAG TGC TGT TGG CTG CAA TCA 270 Phe Thr Gln Gln Leu Leu Gly Cys Leu Gln Cys Cys Trp Leu Gln Ser -10 GGC AGA GCC CCA GCT ACC TAT TAC CTT GTG GAG AGT ATT GAA AAG TCA 318 Gly Arg Ala Pro Ala Thr Tyr Tyr Leu Val Glu Ser Ile Glu Lys Ser GCA CAT GGC TCT GTA TTA NGT ACT TAT GAT CAA ACT CAG ACT CGC ATA 366 Ala His Gly Ser Val Leu Xaa Thr Tyr Asp Gln Thr Gln Thr Arg Ile GGC AGG 372

(2) INFORMATION FOR SEQ ID NO: 167:

Gly Arg

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 343 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 158..337
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.7 seq XTCASXNPSQCLA/AF

(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 167:	
ACAGAATCTT	TAGGTGGGCC TGTTGGTGAG GTCACTTTTC CCTAATGGTA TATTCCAGTT	60
CCTGTAGATC	CTATTCCAGT TCCCAGGACA TATTCCAACC TCGACCTCCA GCCAACTTTG	120
AACCCCTGAA	GTTGTGTGCT GATGTGTTTC TAACAAC ATG GTC TCA CCC AAA GAT Met Val Ser Pro Lys Asp -60 -55	175
CTT CCT CTT Leu Pro Leu	T GTG CTT TTG CAG GAC ATT AAA GTT CCC AGC TCC ATG ACT u Val Leu Leu Gln Asp Ile Lys Val Pro Ser Ser Met Thr -50 -45 -40	223
GGA TCA CAT Gly Ser His	T GCT GGA AAC CCT CAT ATA GAA AGG AAT GAT CTC CCC AGA s Ala Gly Asn Pro His Ile Glu Arg Asn Asp Leu Pro Arg -35 -30 -25	271
CAT GGT TCT His Gly Ser -20	r Pro Gin Phe Phe Thr Gly Xaa Thr Cys Ala Ser Xaa Asn	319
CCA TCT CAG Pro Ser Gln -5	G TGT CTG GCA GCA TTT n Cys Leu Ala Ala Phe 1	343
(i) S	ATION FOR SEQ ID NO: 168: SEQUENCE CHARACTERISTICS: (A) LENGTH: 78 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR MOLECULE TYPE: CDNA ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens	
(ix) 1	(F) TISSUE TYPE: Prostate FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 145 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.6 seq FXSLFCLYFSCFL/HI	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 168:

(2) INFORMATION FOR SEQ ID NO: 169:

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

	()	i) SI	(A) (B) (C)	NCE (LENC TYPE STRA	TH: : NU ANDEC	207 ICLEI NESS	base C AC S: DC	e pai CID DUBLE							
	(j	ii) N	MOLE	CULE	TYPE	E: CI	ONA								
	7 }	/i) ((A)	NAL ORGA TISS	NISM	1: Hc		-		prost	ate				
	·		(B) (C) (D)	JRE: NAME LOCA IDEN OTHE	TION TIFI R IN	: 10 CATI FORM	O14 ON N	14 1ETHC DN:	D: \ scor seq	e 3. ALLE	6 CLIDS	PECI			
ACTO	GGA <i>l</i>	Me					ne G					lu Le		rg TT <i>F</i> aa Leu	
				TAC Tyr											99
				TTG Leu											147
				TTA Leu											195
		ACC Thr 20													207
(2)	INF	orma'	TION	FOR	SEQ	ID I	10:	170:							
	(:	i) SI	(A) (B) (C)	NCE (LEN(TYPI STRA TOP(STH: E: NU ANDE	418 JCLEI ONESS	base [C AC S: DC	e pai CID DUBLE							
	(:	ii) l	MOLE	CULE	TYP	E: CI	ONA								

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П	 х і	 r	۹.	u	Κ.	•. '	-

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 299..379
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 3.6

seq LTLLLITPSPSPL/LF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 170:

ACCTTGGGCT CCAAATTCTA GCTCATAAAG ATGCAAGTKT TGCAATTTCC TATAAATGGT TAAGAAAAGA GCAAGCTGTC CAGAGAGTGA GAAGTTTGAA AAGAGAGGTG CATAAGAGAG 120 AAATGATGTC CATTTGAGCC CCACCACGGA GGTTATGTGG TCCCAAAAGG AATGATGGCC 180 AAGCAATTAA TTTTTCCTCC TAGTTCTTAG CTTGCTTCTG CATTGATTGG CTTTACACAA 240 CTGGCATTTA GTCTGCATTA CACAAATAGA CACTAATTTA TTTGGAACAA GCAGCAAA 298 ATG AGA ACT TTA TTT GGT GCA GTC AGG GCT CCA TTT AGT TCC CTC ACT Met Arg Thr Leu Phe Gly Ala Val Arg Ala Pro Phe Ser Ser Leu Thr -20 CTG CTT CTA ATC ACC CCT TCT CCC AGC CCT CTT CTA TTT GAT AGA GGT 394 Leu Leu Ile Thr Pro Ser Pro Ser Pro Leu Leu Phe Asp Arg Gly CTG TCC CTC AGA TCA GCA ATG TCG 418 Leu Ser Leu Arg Ser Ala Met Ser 10

(2) INFORMATION FOR SEQ ID NO: 171:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 238 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: 107..229
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.6

seq AVSSLIAVGTSHG/LA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 171:

AAGGAAGAAG AAATTACCTG ATTCTTTTTC ACTTCATGGA TCAGTT ATG CGC CAT Met Arg His -40	115
TCA CTT TTG AAG GGA ATT TCT GCC CAG ATA GTG TCT GCA GCT GAC AAA Ser Leu Leu Lys Gly Ile Ser Ala Gln Ile Val Ser Ala Ala Asp Lys -35 -30 -25	163
GTA GAT GCT GGC TTG CCT ACA GCA ATT GCA GTA TCC AGT CTG ATA GCA Val Asp Ala Gly Leu Pro Thr Ala Ile Ala Val Ser Ser Leu Ile Ala -20 -15 -10	211
GTG GGT ACA TCT CAT GGA TTG GCT GGG Val Gly Thr Ser His Gly Leu Ala Gly -5 1	238
(2) INFORMATION FOR SEQ ID NO: 172: (i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 188 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 120164 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.5</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 172:	
TGTGAAGATG ACAGAGATCT AACTTCTGAG AGCAGAGGTG TCAAGTGACG GTCCCCTTGG	60
AGGAATGGTC TTTGCATCTG ACTACTTCCT TCTGCAACTG TGTTCTTCCA TTAGCTTCC	119
ATG ACA CTC TCC TGC TTT ATT TTT TTC TAC ATC TCT AGC CTT TGC TGT Met Thr Leu Ser Cys Phe Ile Phe Phe Tyr Ile Ser Ser Leu Cys Cys -15 -5 1	167
TTC CTC TCC TAC CCC ACC AGG Phe Leu Ser Tyr Pro Thr Arg 5	138
(2) INFORMATION FOR SEC ID NO. 173.	

- (2) INFORMATION FOR SEQ ID NO: 173:
 - (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 168 base pairs

(B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 2872 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.5</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 173:	
ATAGATCAGT GACGTCTTTT TCTTCAG ATG ATC CTA TGT TTC CTT CCT CAT Met Ile Leu Cys Phe Leu Leu Pro His -15 -10	54
CAT CGT CTT CAG GAA GCC AGA CAG ATT CAA GTA TTG AAG ATG CTG CCA His Arg Leu Gln Glu Ala Arg Gln Ile Gln Val Leu Lys Met Leu Pro -5 1 5 10	102
AGG GAA AAA TTA AGR AGA AGR AGA AGA GAG AAA ACA AAT AAA TGG GAA Arg Glu Lys Leu Arg Arg Arg Arg Glu Lys Thr Asn Lys Trp Glu 15 20 25	150
AAA AGA AAG GGC AGC GGG Lys Arg Lys Gly Ser Gly 30	168
(2) INFORMATION FOR SEQ ID NO: 174:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 135 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR 	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 64105</pre>	

(C) IDENTIFICATION METHOD: Von Heijne matrix

seq FSLFALNMPLGFC/VY

(D) OTHER INFORMATION: score 3.5

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 174:

TTT	ATTTI	'AA (CCATO	CTTT	ra ci	TATT	rtta(G AAG	GGAA	ACTA	GCT	rtag:	rag	TGGG'	TTGCCC	60
TGT	ATG Met	TTT Phe	TCT Ser	CTT Leu	TTT Phe -10	GCT Ala	CTT Leu	AAT Asn	ATG Met	CCA Pro -5	TTG Leu	GGT Gly	TTT Phe	TGT Cys	GTG Val 1	108
	GTG Val															135
(2)	INFC	RMAT	NOI	FOR	SEQ	ID 8	NO: 1	175:								
	(i	.) SE	(B) (C)	LENG TYPE STRA	CHARA STH: C: NU NDEC OLOGY	303 ICLEI NESS	base C AC S: DC	e pai CID OUBLE								
	(i	i) M	OLEC	CULE	TYPE	: CI	ANC									
	(v	·i) (ORGA	SOUP NISM SUE T	l: Ho		-		rost	ate					
	(i	×) E	(B) (C)	NAME LOCA IDEN	C/KEY ATION ITIFI CR IN	: 16 CATI	32 ON M	255 1ETHC	D: V	e 3.	5	ne ma				
	(x	i) S	SEQUE	NCE	DESC	RIPT	NOI:	SEÇ) ID	NO:	175	:				
ATT'	rgatt	TT A	AGTC	AGGGT	rg TA	\AGA/	ATATO	G TAT	TATI	rgtt	CCC	LAAA	AAA	TCTG:	rgtaa?	k 60
AAC'	TTCAT	'AG 1	rgtgi	AAAC	AG TO	GCA	ACTGS	s KTC	SATTA	AAAA	CAT	CATT	rag	AAAA	GACACT	120
CTT	CCTG	STT T	rtga <i>i</i>	AATT(GA CT	CCT	CAAA	A GG <i>i</i>	ACAGO	CTGA		Met A		TCT :		174
	GGT Gly															222
	GGC Gly -10															270
	GCC Ala															303

•	VO 99	9/0655	50				13	134								
	(.	i) S	(A) (B) (C)	LENG TYPI STR	CHAR GTH: E: NI ANDEI OLOGY	317 JCLE: ONES:	bas IC A S: D	e pa: CID OUBL								
	(:	ii) l	MOLE	CULE	TYPI	E: C	DNA									
	(,	vi) ((A)	ORG	SOUI ANISM SUE 1	4: Ho										
			(B) (C) (D)	NAMI LOCA IDEN	E/KEY ATION NTIFI ER IN	1: 60 [CAT] IFORM	O19 ION N	94 METHO ON:	DD: N scor seq	re 3. LPRI	.5 LLSLS	SQHSI				
	,,	, .	o no go i	311,013	0000	MIT.	TION	. SE(עני ע	NO:	1/6	•				
AGA	GTTT(CCG (GTCT	GGC'	TT TO	GCG	GGTC	r gg	rttg/	AAGC	TCT	CCTG	TTT	GACG	AAAGT	59
ATG Met -45	TCT Ser	CAG Gln	GAA. Glu	GGT Gly	GCG Ala -40	GTC Val	CCA Pro	GCT Ala	AGC Ser	GCG Ala -35	GTT Val	CCC Pro	CTG Leu	GAA Glu	GAA Glu -30	107
TTA Leu	AGT Ser	AGC Ser	TGG Trp	CCA Pro -25	GAG Glu	GAG Glu	CTA Leu	TGC Cys	CGC Arg -20	CGG Arg	GAA Glu	CTG Leu	CCG Pro	TCC Ser -15	GTC Val	155
CTG Leu	CCC Pro	CGA Arg	CTC Leu -10	CTC Leu	TCA Ser	TTG Leu	TCT Ser	CAA Gln -5	CAT His	TCT Ser	GAA Glu	AGT Ser	TGG Trp 1	ATT Ile	GAG Glu	203
CAT His	ATT Ile 5	CAA Gln	ATT Ile	TTG Leu	AAA Lys	ATT Ile 10	ATT Ile	GTA Val	GAA Glu	ATG Met	TTT Phe 15	TTA Leu	CCT Pro	CAT His	ATG Met	251
AAC Asn 20	CAC His	CTG Leu	ACA Thr	TTG Leu	GAA Glu 25	CAG Gln	ACT Thr	TTC Phe	TTT Phe	TCA Ser 30	CAA Gln	GTG Val	TTA Leu	CCA Pro	AAG Lys 35	299

317

(2) INFORMATION FOR SEQ ID NO: 177:

ACT GTG AAA TTA TTC GAT

Thr Val Lys Leu Phe Asp

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 370 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:

(ix)

(xi)

50	135	PCT/IB98/0
	ORGANISM: Homo Sapiens TISSUE TYPE: Cancerous prostate	
FEAT	URE:	
(A)	NAME/KEY: sig_peptide	
	LOCATION: 254361	
(C)	IDENTIFICATION METHOD: Von Heijne matrix	
	OTHER INFORMATION: score 3.5	
	seq AAVVFAVVLSIHA/TV	
SEQUI	ENCE DESCRIPTION: SEQ ID NO: 177:	
AGGA	AGGCTG CAGAGTGGCG ACGTCTACGC CGTAGGTTGG AGGCTGT	GGG 60

AGTAACTGTG GGGTGGCCGG GCGCCAGCTC CCAGGCCGCA GAAGTGACCT GCGGTGGAGT TCCCTCCTCG 120 CTGCTGGAGA ACGGAAGGGA ARAAGGTTSC TGGCCGGGTG AAAGTGCCTC CCTCTGCTTG 180 ACGGGGCTGA GGGGCCCGAA GTCTAGGGCG TCCGTAGTCG CCCCGGCCTC CGTGAAGCCC 240 CAGGTCTAGA GAT ATG ACC CGA GAG TGC CCA TCT CCG GCC CCG GGG CCT 289 Met Thr Arg Glu Cys Pro Ser Pro Ala Pro Gly Pro -35 -30 GGG GCT CCG CTG AGT GGA TCG GTG CTG GCA GAG GCG GCA GTA GTG TTT 337 Gly Ala Pro Leu Ser Gly Ser Val Leu Ala Glu Ala Ala Val Val Phe -15 GCA GTG GTG CTG AGC ATC CAC GCA ACC GTA TGG 370 Ala Val Val Leu Ser Ile His Ala Thr Val Trp -5

(2) INFORMATION FOR SEQ ID NO: 178:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 470 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 369..470
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 92 region 2..103 id AA059664 est
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide (B) LOCATION: 216..269

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 14.8
seq LLWWALLLGLAQA/CP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 178:

AAGTGGATGG TTCCAGGCAC CCTGTCTGGG GCAGGGAGGG CACAGGCCTG CACATCGAAG 60 GTGGGGTGGG ACCAGGCTGC CCCTCGCCCC AGCATCCAAG TCCTCCCTTG GGCGCCCGTG GCCCTGGCAG ACTCTCAGGG CTAAGGTCCT CTGTTGCTTT TTGGTTCCAC CTTAGAAGAG GCTCGCTTGA CTAAGAGTAG CTTGAAGGAG GCACC ATG CAG GAG CTG CAT CTG 233 Met Gln Glu Leu His Leu -15 CTC TGG TGG GCG CTT CTC CTG GGC CTG GCT CAG GCC TGC CCT GAG CCC 281 Leu Trp Trp Ala Leu Leu Gly Leu Ala Gln Ala Cys Pro Glu Pro -10 1 TGC GAC TGT GGG GAA AAG TAT GGC TTC CAG ATC GCC GAC TGT GCC TAC 329 Cys Asp Cys Gly Glu Lys Tyr Gly Phe Gln Ile Ala Asp Cys Ala Tyr 10 CGC GAC CTA GAA TCC GTG CCG CCT GGC TTC CCG GCC AAT GTG ACT ACA Arg Asp Leu Glu Ser Val Pro Pro Gly Phe Pro Ala Asn Val Thr Thr 25 30 CTG AGC CTG TCA GCC AAC CGG CTG CCA GGC TTG CCG GAR GGT GCC TTC 425 Leu Ser Leu Ser Ala Asn Arg Leu Pro Gly Leu Pro Glu Gly Ala Phe 40 AGG GAG GTG CCC CTG CTG CAG TCG CTG TGG CTG GCA CAC AAT GAG 470 Arg Glu Val Pro Leu Leu Gln Ser Leu Trp Leu Ala His Asn Glu 60

(2) INFORMATION FOR SEQ ID NO: 179:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 331 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 69..328
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95 region 1..260 id H96534 est

WO 99/06550	137	PCT/IB98/01232

(i	x) F	ΈA	T	JR	Ε:
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- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 14..67
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 13.6

seq LLLLALCATGAQG/LY

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 179:

CTCT	rctg(CGG (GGG (Gly <i>P</i>	Arg (Leu 1	-		 49
				GGG Gly							97
				GAA Glu							145
				TGG Trp							193
				ATG Met							241
				CAG Gln							289
				GAC Asp 80			 				331

(2) INFORMATION FOR SEQ ID NO: 180:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 195 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 90..129
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 1..40

138

id AA134726 est

1	lix) F	F.A	TH	RE:

- (A) NAME/KEY: other
- (B) LOCATION: 157..195
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 66..104 id AA134726

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 124..156
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 93

region 34..66 id AA134726

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 107..195
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 1..89 id R17226

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 76..138
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 12.7

seq ILFLLSWSGPLQG/QQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 180:

AAGCTAACCC TCGGGCTTGA GGGGAAGAGG CTGACTGTAC GTTCCTTCTA CTCTGGCACC

ACTCTCCAGG CTGCC ATG GGG CCC AGC ACC CCT CTC CTC ATC TTG TTC CTT

Met Gly Pro Ser Thr Pro Leu Leu Ile Leu Phe Leu

-20

-15

TTG TCA TGG TCG GGA CCC CTC CAA GGA CAG CAG CAC CAC CTT GTG GAG
Leu Ser Trp Ser Gly Pro Leu Gln Gly Gln Gln His His Leu Val Glu

-5

TAC ATG GAA CGC CGA CTA GCT GCT TTA GAG GAA CGG

Tyr Met Glu Arg Arg Leu Ala Ala Leu Glu Glu Arg

10

15

(2) INFORMATION FOR SEQ ID NO: 181:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 352 base pairs

(B) TYPE: NUCLEIC ACID

MO 33/00220				139												rC1/1D70	
				STRA					:								
	(i:	i) M	OLEC	CULE	TYPE	e: cr	ANG										
	(v:	i) C	(A)	NAL ORGA TISS	NISM	1: Hc		-									
	(i:	×) F	(B) (C)	JRE: NAME LOCA IDEN OTHE	TION	: 31 :CATI	33 ON M	1ETHC	DD: b iden regi id T est	tity on 7	, 97 43	3					
	(i:	×) F	(B) (C)	JRE: NAME LOCA IDEN OTHE	TION TIFI	: 11 CATI	.91 ON M	.99 IETHO	D: V	e 8.	8		tri;				
	(x.	i) S	EQUE	ENCE	DESC	CRIPT	CION:	SEÇ	Q ID	NO:	181:	:					
ACGT	TACC'	TT T	rggg:	rggto	GG T	rttc/	ATTC	TG:	rgcco	SCCT	GCT'	rctgo	GGC (CAGTO	GATCCA	60	
GGTGTCTGGT GACC				ACCC	GG GG	CACAC	GCTG	C TTO	GCT	SCTG	TGG	GCAC	CTC A	AGCT	rccc	118	
	TCC Ser															166	
	CTG Leu															214	
	CTG Leu															262	
	TCT Ser															310	
	TCA Ser															352	
(2)	INFO		EQUEI (A) (B)	FOR NCE (LEN(TYP! STRA	CHAR GTH: E: NO	ACTE 447 UCLE:	RIST base	ICS: e pa: CID									

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 113..306
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 71..264 id H83784

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 42..111
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 1..70

id H83784

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 378..414
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 346..382

id H83784

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 305..340
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 264..299

id H83784

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 250..350
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 2..102

id W32197

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (3) LOCATION: 392..449
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 142..199

id W32197

est

	(1)	i x) - 1	(B) (C)	URE: NAME LOCA IDEN OTHE	ATION NTIF	N: 34	193 [ON N	4ETHC	ider regi	olast ntity ion 1 V3219	7 100 100.					
	()	x) l	(B) (C)	JRE: NAME LOCA IDEN OTHE	ATION NTIFI	1: 39 [CAT]	74 ON N	1ETHC	ider regi	olast htity on 1 13725	/ 100 L53					
	(i	x) I	(B) (C)	JRE: NAME LOCA I DEN OTHE	TION TIFI	1: 85 CAT	015 ON M	50 1ETHC	D: V	e 8.	5		atri: CG/DE			
	()	(i) S	SEQUE	ENCE	DESC	CRIPT	CION:	SEQ) ID	NO:	182:					
AACI	rtgto	STC (CGGGT	rggwi	RG AC	CTGG!	ATTA	G CTO	GCGG/	ASCC	TGG	AAGC:	rgc (CTGT	CCTTCT	60
CCCI	rgtgo	CTT A	AACC	AGAGO	GT GO			Sly 1					Leu '	STC A Val 1		111
GCA Ala	GCA Ala	CTG Leu	TTA Leu -10	CTG Leu	GGT Gly	CTC Leu	ATG Met	ATG Met -5	GTG Val	GTC Val	ACT Thr	GGA Gly	GAC Asp 1	GAG Glu	GAT Asp	159
GAG Glu	AAC Asn 5	AGC Ser	CCG Pro	TGT Cys	GCC Ala	CAT His 10	GAG Glu	GCC Ala	CTC Leu	TTG Leu	GAC Asp 15	GAG Glu	GAC Asp	ACC Thr	CTC Leu	207
TTT Phe 20	TGC Cys	CAG Gln	GGC Gly	CTT Leu	GAA Glu 25	GTT Val	TTC Phe	TAC Tyr	CCA Pro	GAG Glu 30	TTG Leu	GGG Gly	AAC Asn	ATT Ile	GGC Gly 35	255
			GTT Val													303
TGG Trp	ATG Met	GAG Glu	CCG Pro 55	ATA Ile	GTC Val	AAG Lys	TTC Phe	CCG Pro 60	GGG Gly	GCC Ala	GTG Val	GAC Asp	GGC Gly 65	GCA Ala	ACC Thr	351
TAT Tyr	ATC Ile	CTG Leu 70	GTG Val	ATG Met	GTG Val	GAT Asp	CCA Pro ·75	GAT Asp	GCC Ala	CCT Pro	AGC Ser	AGA Arg 80	GCA Ala	GAA Glu	CCC Pro	399
AGA Arg	CAG Gln 85	AGA Arg	TTC Phe	TGG Trp	AGA Arg	CAT His	TGG Trp	CTG Leu	GTA Val	ACA Thr	GAT Asp	ATC Ile	AAG Lys	GGC Gly	GCC Ala	447

(2) INFORMATION FOR SEQ ID NO: 183:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 217 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 125..182
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 6..63 id R18560

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 176..213
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 92

region 58..95 id R18560

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 145..182
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100

region 1..38 id R13864

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 176..213
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 92

region 33..70

id R13864

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 176..213
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 92

region 2..39

id HSC01E071

est

(ATURE: A) NAME/KEY: sig_peptide B) LOCATION: 119190 C) IDENTIFICATION METHOD: Von Heijne matrix D) OTHER INFORMATION: score 7.3 seq VHLLSLCSGKVYA/RM
(xi) SE	QUENCE DESCRIPTION: SEQ ID NO: 183:
ACTGGGAGCC GC	CTCCGTCG CCGCCGTCAG AGCCGCCCTA TCAGAGTTCC TACCANTTTG 60
TGGTTCCAGC AG	CTTCTGTT CCAGATTATC TTAACAAGAA AACCAACTGG AAAAAAAA 118
	TT ATC TTC GCA TTT TTC GGT GGT GTT CAC CTT TTA TCC eu Ile Phe Ala Phe Phe Gly Gly Val His Leu Leu Ser -20 -15 -10
	GG AAA GTA TAT GCA AGA ATG GCA TCT CTA AGA GGA CTC 214 ly Lys Val Tyr Ala Arg Met Ala Ser Leu Arg Gly Leu -5 5
GGG Gly	217
(2) INFORMATI	ON FOR SEQ ID NO: 184:
(UENCE CHARACTERISTICS: A) LENGTH: 433 base pairs B) TYPE: NUCLEIC ACID C) STRANDEDNESS: DOUBLE D) TOPOLOGY: LINEAR
(ii) MO	LECULE TYPE: CDNA
(IGINAL SOURCE: A) ORGANISM: Homo Sapiens F) TISSUE TYPE: Cancerous prostate
(ATURE: A) NAME/KEY: other B) LOCATION: 139361 C) IDENTIFICATION METHOD: blastn D) OTHER INFORMATION: identity 99 region 92314 id AA100852 est
(ATURE: A) NAME/KEY: other B) LOCATION: 360434 C) IDENTIFICATION METHOD: blastn D) OTHER INFORMATION: identity 100 region 314388 id AA100852 est

(ix) FEATURE:

WO 99/06550 144 (A) NAME/KEY: other (B) LOCATION: 139..434 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 100..395 id AA224847 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 139..361 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 92..314 id AA161042 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 368..434 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 323..389 id AA161042 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 139..365 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 87..313 id H64488 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 52..144 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 92 region 1..93 id H64488 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 171..396 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 129..354 id AA088770 est (ix) FEATURE: (A) NAME/KEY: sig_peptide

(B) LOCATION: 167..253

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 7.1

seq LIFLCGAALLAVG/IW

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 184:

AAA	AGC	SCC '	racco	CTGC	CT GO	CAGG	rgago	AG1	rggto	STGT	GAG	AGCC	AGG (CGTC	сстстс	G 60
CCTC	CCC/	ACT (CAGTO	GCA	AC AC	CCCG	GGAG	TG	ידדדמ	STCC	TTTC	STGG	AGC (CTCA	GCAGT	r 120
CCCT	CTT	rca (GAACT	TYRV	rk Go	CCAA	GAGCO	СТО	GAACA	AGGA	GCCA			CAG : Gln (175
			ATT Ile													223
			GCA Ala													271
			TTT Phe 10													319
			AAC Asn													367
			GGT Gly													415
			GTG Val													433

(2) INFORMATION FOR SEQ ID NO: 185:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 372 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 128..242
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 92 region 1..115 id R58075 est
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 (B) LOCATION: 220..303

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 6.6

seq IVSLLGFVATVTL/IP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 185:

AAGATAGGCG GGTGCAGCGG GGCAGAACAT AGGTTGCCTT AGAGAGGTTC CCCGGAGTCC 60 CGACGGCGGC TCAAGTCAGA GTTGCTGGGT TTTGCTCAGA TTGGTGTGGG AAGAGCCTGC 120 CTGTGGGGAG CGGCCACTCC ATACTGCTGA GGCCTCAGGA CTGCTGCTCA GCTTGCCCGT TACCTGAAGA GGCGGCGGAS GGGCCCCTGA CCGGTCACC ATG TGG GCC TTC TCG 234 Met Trp Ala Phe Ser GAA TTG CCC ATG CCG CTG CTG ATC AAT TTG ATC GTC TCG CTG GGA 282 Glu Leu Pro Met Pro Leu Leu Ile Asn Leu Ile Val Ser Leu Leu Gly TTT GTG GCC ACA GTC ACC CTC ATC CCG GCC TTC CGG GGC CAC TTC ATT 330 Phe Val Ala Thr Val Thr Leu Ile Pro Ala Phe Arg Gly His Phe Ile -5 GCT GCG CGC CTC TGT GGT CAG GAC CTC AAC AAA ACC AGC CAG 372 Ala Ala Arg Leu Cys Gly Gln Asp Leu Asn Lys Thr Ser Gln 15

(2) INFORMATION FOR SEQ ID NO: 186:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 402 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 112..403
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 33..324 id H97426

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 59..295
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 2..239 id W44834

est

(ix) FEATURE: (A) NAME/KEY: other

(B) LOCATION: 106..156

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 4..54 id R57989

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 161..190

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93

region 62..91 id R57989

est

(ix) FEATURE:

(A) NAME/KEY: sig peptide

(B) LOCATION: 148..204

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 6.3

seq VLMRLVASAYSIA/QK

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 186:

AGCTGAGGTA GGGATGCSAT CCTTCTCAAA AGACTTATTG ACAGTGCCAA AGCTSGGTAC

TGGACACAAC GAGGGACCTG GGTCTACGAT AACGCGCTTK TGCTCCTCCT GAAGTGTCTT

TGGTCCAACG TTGTTCCAGA GTGTACC ATG GCT TCC AGT AAC ACT GTG TTG ATG 174

Met Ala Ser Ser Asn Thr Val Leu Met

-15

CGG TTG GTA GCC TCC GCA TAT TCT ATT GCT CAA AAG GCA GGD ATG ATA 222 Arg Leu Val Ala Ser Ala Tyr Ser Ile Ala Gln Lys Ala Gly Met Ile

-10 -5 1

GTC AGA CGT GTT ATT GCT GAA GGA GAC CTG GGT ATT GTG GAG ADG ACC

Val Arg Arg Val Ile Ala Glu Gly Asp Leu Gly Ile Val Glu Xaa Thr

15 20

TGT GCA ACA GAC CTG CAG ACC AAA GCT GAC CGA TTG GCA CAG ATG AGN 318 Cys Ala Thr Asp Leu Gln Thr Lys Ala Asp Arg Leu Ala Gln Met Xaa

> 25 . 30

ATA TGT TCT TCA TTG GCC CGG AAA TTC CCC AAA CTC ACA ATT ATA GGG

Ile Cys Ser Ser Leu Ala Arg Lys Phe Pro Lys Leu Thr Ile Ile Gly 40

45

GAA GAG GAT CTG CCT TCT RMG GAA GTG GAT CAA GAG 402

Glu Glu Asp Leu Pro Ser Xaa Glu Val Asp Gln Glu

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 317 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 111..318
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 6..213

id R18560

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 131..318
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 1..188

id R13864

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 162..318
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 1..157

id HSC01E071

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 207..318
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 1..112

id AA016124

est

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (3) LOCATION: 105..176
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq VHLLSLCSGKAIC/KN

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 187:

Met Lys Phe Leu														116	
				TTC Phe											164
				AAG Lys 1											212
				GCC Ala											260
				GGT Gly											308
	CTG Leu														317

(2) INFORMATION FOR SEQ ID NO: 188:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 499 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 160..401
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95

region 59..300

id H29377

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 454..499
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100

region 356..401

id H29377

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 136..179
 - (C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 36..79 id H29377

est

(ix) FEATURE:

- (A) NAME/KEY: other(B) LOCATION: 397..436
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 297..336

id H29377

est

(ix) FEATURE:

- (A) NAME/KEY: other(B) LOCATION: 135..295
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 95

region 293..453

id N28905

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 45..127
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 93

region 4..86

id N28905

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 334..388
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 489..543

id N28905

est

-(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 135..395
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 81..341

id H11885

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 160..384
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 95

region 105..329

id H15231

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 136..181

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93

region 82..127 id H15231 est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 146..298

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.9

seq ALXVLPLLGLHEA/AS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 188:

AACI	TCCC	GG '	TTCG	GCAAT	ra ao	CCTGC	SAGCO	GGG	CGGCC	STAG	GTT	GCTC	TT 1	raggo	GCTTCA	60
ccc	GAAC	SCT (CCAC	CTTCC	SC TO	CCCG1	CTTT	CTC	GAAA	ACAC	CGC1	TTGA	ATC :	rcggo	CGGTGC	120
GGG	ACAGA	ACG (CTAGT	rgtg <i>i</i>	AG CO	CNMC								GGC Gly		172
			GGC Gly													220
			CTT Leu													268
			CTT Leu													316
			GCA Ala 10													364
			TTC Phe													412
GAG Glu	GAC Asp 40	AGC Ser	TGC Cys	CAC His	TAC Tyr	CTG Leu 45	TTG Leu	TAT Tyr	TCA Ser	CTC Leu	ATC Ile 50	TTT Phe	GTA Val	AAT Asn	TCC Ser	460
			ACA Thr													499

(2) INFORMATION FOR SEQ ID NO: 189:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 219 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

WO 99/06550 PCT/IB98/01232

	(D)	TOPOLOGY	: LINE	AR								
(i:) MOLE	CULE TYPE	E: CDNA									
(v :	(A)	GINAL SOUF ORGANISM TISSUE T	1: Homo	Sapi ancer	ens ous	pros	state	è				
(i)	(B) (C)	TURE: NAME/KEY LOCATION IDENTIFI OTHER IN	: 452 CATION	21 METH	ide reg	ntit ion HUMH	stn Sy 93 11 BC46	.77				
(ix	(B) (C)	URE: NAME/KEY LOCATION IDENTIFIC OTHER IN	: 632 CATION	21 METH	ide reg	ntit ion	tn y 94 11 0569	59				
(ix	(B) (C)	URE: NAME/KEY LOCATION IDENTIFIC OTHER INI	: 124	METHO	ide: reg:	ntity	y 97 97	132				
	(B) (C) (D)	JRE: NAME/KEY: LOCATION: IDENTIFIC OTHER INE	: 172 CATION N	METHO ON:	D: V scor seq	e 5. XVL	. 5 /LSV\	/XXA				
	_	7		. 000	2 10	NO.	109	•				
ATG CGT T Met Arg Pl	CC CGC ne Arg	CAT TTT 1 His Phe 2 -20	CGM AAA Kaa Lys	TWA Xaa	ATT Ile -15	GGG Gly	MAG Xaa	GTA Val	CTG Leu	GTT Val -10	TTA Leu	48
AGT GTA G Ser Val Va	T SCC 1 Xaa -5	GMC GCA A Xaa Ala N	ATG GCA Met Ala	GCC Ala 1	TTT Phe	GCA Ala	GTG Val	SHA Xaa 5	CCT Pro	CAG Gln	GGG Gly	96
CCC GCG T1 Pro Ala Le 10	A SSM u Xaa	TCT GAA C Ser Glu F	CCA MTG Pro Xaa 15	MTG Xaa	CYG Xaa	GGT Gly	TCA Ser 20	CCC Pro	ACA Thr	TCT Ser	CCA Pro	144
AAG CCA GC Lys Pro G1 25	SA GTT y Val	AAT GCC C Asn Ala G 30	CAG TTC	TTA Leu	CCT Pro	GGA Gly 35	TTT Phe	TTA Leu	ATG Met	GGG Gly	GMT Xaa 40	192

TTG CCA GCT CCG GTG ACT CCA CAA CCT Leu Pro Ala Pro Val Thr Pro Gln Pro 45

219

(2) INFORMATION FOR SEQ ID NO: 190:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 483 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 105..414
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 1..310 id T26956

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 45..359
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 1..315 id T31666

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 202..332
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 137..267

id R14990

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 127..201
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 63..137

id R14990

est

(ix) FEATURE:

- (A) NAME/KEY: other
 (B) LOCATION: 65..114
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 1..50 id R14990 est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide(B) LOCATION: 1..120

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.2

seq LCVEFASVASCDA/AV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 190:

ATG Met -40	GIU	TTG Leu	GGG Gly	AGT Ser	TGC Cys -35	CTG Leu	GAG Glu	GGC Gly	GGG Gly	AGG Arg -30	Glu	GCG Ala	GCG Ala	GAG Glu	GAA Glu -25	48
GAG Glu	GGC Gly	GAG Glu	CCT Pro	GAG Glu -20	GTG Val	AAA Lys	AAG Lys	CGG Arg	CGA Arg -15	CTT Leu	CTG Leu	TGT Cys	GTG Val	GAG Glu -10	TTT Phe	96
GCC Ala	TCG Ser	GTC Val	GCA Ala -5	AGC Ser	TGC Cys	GAT Asp	GCC Ala	GCA Ala 1	GTG Val	GCT Ala	CAG Gln	TGC Cys 5	TTC Phe	CTG Leu	GCC Ala	144
GAG Glu	AAC Asn 10	GAC Asp	TGG Trp	GAG Glu	ATG Met	GAA Glu 15	AGG Arg	GCT Ala	CTG Leu	AAC Asn	TCC Ser 20	TAC Tyr	TTC Phe	GAG Glu	CCT Pro	192
CCG Pro 25	GTG Val	GAG Glu	GAG Glu	AGC Ser	GCC Ala 30	TTG Leu	GAA Glu	CGC Arg	CGA Arg	CCT Pro 35	GAA Glu	ACC Thr	ATC Ile	TCT Ser	GAG Glu 40	240
CCC	AAG Lys	ACC Thr	TAT Tyr	GTT Val 45	GAC Asp	CTA Leu	ACC Thr	AAT Asn	GAA Glu 50	GAA Glu	ACA Thr	ACT Thr	GAT Asp	TCC Ser 55	ACC Thr	288
ACT Thr	TCT Ser	AAA Lys	ATC Ile 60	AGC Ser	CCA Pro	TCT Ser	GAA Glu	GAT Asp 65	ACT Thr	CAG Gln	CAA Gln	GAA Glu	AAT Asn 70	GGC Gly	AGC Ser	336
ATG Met	TTC Phe	TCT Ser 75	CTC Leu	ATT Ile	ACC Thr	TGG Trp	AAT Asn 80	ATT Ile	GAT Asp	GGA Gly	TTA Leu	GAT Asp 85	CTA Leu	AAC Asn	AAT Asn	384
CTG Leu	TCA Ser 90	GAG Glu	AGG Arg	GCT Ala	CGA Arg	GGG Gly 95	GTG Val	TGT Cys	TCC Ser	TAC Tyr	TTA Leu 100	GCT Ala	TTG Leu	TAC Tyr	AGC Ser	432
CCA Pro 105	GAT Asp	GTG Val	ATA Ile	TTT Phe	CTA Leu 110	CAG Gln	GAA Glu	GTT Val	ATT Ile	CCC Pro 115	CCA Pro	TAT Tyr	TAT Tyr	AGC Ser	TAC Tyr 120	480
CTA Leu																483

⁽²⁾ INFORMATION FOR SEQ ID NO: 191:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 444 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Hypertrophic prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 182..401

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 165..384

id W56608

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 45..130

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 30..115 id W56608

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 127..191

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 111..175

id W56608

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 401..446

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 385..430

id W56608

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 311..446

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 1..136

id R17248

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(3) LOCATION: 13..378

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5

seq RLVVVSVSPQSRA/SL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 191:

AGT	GCGG	CCG	TC A	TG G	ia s	CG C er P 120	CC T	TC A	GC G er G	ly A	CG C la L	TG C eu G	AG C	TG A	CG GAC hr Asp -110	51
CTG Leu	GAT Asp	GAC Asp	TTC Phe	ATC Ile -10	GLy	CCG Pro	TCT Ser	CAG Gln	GAG Glu -10	Cys	ATC Ile	AAG Lys	CCT Pro	GTC Val -95	AAA Lys	99
GTG Val	GAA Glu	AAA Lys	AGG Arg -90	GCG Ala	GGA Gly	AGT Ser	GGC Gly	GTG Val -85	GCC Ala	AAG Lys	ATT Ile	CGC Arg	ATT Ile -80	GAA Glu	GAT Asp	147
GAC Asp	GGG Gly	AGC Ser -75	TAC Tyr	TTC Phe	CAA Gln	ATT Ile	AAC Asn -70	CAA Gln	GAC Asp	GGC Gly	DGG Xaa	ACC Thr -65	CGG Arg	AGG Arg	CTG Leu	195
GAG Glu	AAG Lys -60	GCC Ala	AAG Lys	GTC Val	TCG Ser	CTA Leu -55	AAC Asn	TAC Tyr	TGC Cys	NWG Xaa	GCG Ala -50	TGC Cys	AGC Ser	GGC Gly	TGC Cys	243
ATC Ile -45	ACC Thr	TCC Ser	GCA Ala	GAG Glu	ACC Thr -40	GTG Val	CTT Leu	ATC Ile	ACC Thr	CAG Gln -35	CAG Gln	AGC Ser	CAC His	GAG Glu	GAG Glu -30	291
CTG Leu	AAG Lys	AAG Lys	GTT Val	CTA Leu -25	GAT Asp	GCT Ala	AAC Asn	AAG Lys	ATG Met -20	GCG Ala	GCA Ala	CCC Pro	AGT Ser	CAG Gln -15	CAG Gln	339
AGG Arg	CTG Leu	GTT Val	GTA Val -10	GTT Val	TCG Ser	GTC Val	TCA Ser	CCA Pro -5	CAG Gln	TCT Ser	AGA Arg	GCA Ala	TCG Ser 1	CTG Leu	GCT Ala	387
GCA Ala	CGG Arg 5	TTT Phe	CAG Gln	CTG Leu	AAW Xaa	CCT Pro 10	ACA Thr	GAT Asp	ACT Thr	GCC Ala	AGG Arg 15	AAA Lys	TTA Leu	ACC Thr	TCA Ser	435
TTC Phe 20																444

(2) INFORMATION FOR SEQ ID NO: 192:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 335 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate

	(:	ix)	(B) (C)	NAM!	ATION VTIF	1: 4: [CAT]	49 ⁻ ION 1	METH	reg	ntity	y 92 L5	1				
	(:	ix)	(B) (C)	NAM!	ATION NTIFI	1: 84 CAT	12: [ON N	15 METHO	DD: \	ce 4.						
	(:	ki) :	SEQUI	ENCE	DESC	CRIP	rion:	: SE(Q ID	NO:	192	•				
ATA	CTTT	CTG	YAGY	AGTC	CT GO	GACC'	CCT	G TG	CAAG	AACA	TGA	AACA	CCT	GTGG	TTCAT	C 60
CTC	CTGC'	r G G	TGGC	AGCT	CC CA					al P					CA GGG la Gl; -3	У
GCT Ala	GGC Gly	TCA Ser	CGA Arg	CTG Leu -30	GTA Val	AAG Lys	CCC Pro	TCA Ser	CAG Gln -25	ACC Thr	CTG Leu	TCC Ser	CTC Leu	ACC Thr -20	TGC Cys	161
GCT Ala	GTC Val	TCT Ser	GGT Gly -15	GGC Gly	TCA Ser	TTA Leu	GTA Val	GCG Ala -10	GAA Glu	CTT Leu	CTT Leu	CTT Leu	GGA Gly -5	GCT Ala	GGA Gly	209
TCC Ser	GGC Gly	AGT Ser 1	CAC His	CTG Leu	GGA Gly	CGG Arg 5	GCC Ala	TGG Trp	AGT Ser	GGA Gly	TTG Leu 10	GGT Gly	TCA Ser	TCT Ser	ATT Ile	257
ATA Ile 15	GAG Glu	GCA Ala	ATA Ile	GTG Val	GGA Gly 20	GTA Val	CTT Leu	CTT Leu	ACA Thr	ATC Ile 25	CGT Arg	CCC Pro	TCA Ser	AGA Arg	CTC Leu 30	305
		Pro	TAT Tyr	His	Trp	Thr										335
(2)			(B) (C)		CHARA STH: C: NU	CTEI 391 ICLEI	RISTI base IC AC S: DC	ICS: e pai CID DUBLE								
			MOLEC ORIGI (A)	NAL ORG <i>i</i>	SOUF MSINA	RCE: 1: Ho	omo S	Sapie	ens							
			(1)	1155	OL I	TPE:	Nor	cmal	pros	state	•					

WO 99/06550 PCT/IB98/01232

(:	ix)	(A) (B) (C)	LOC	ME/KI CATIO ENTII	ON: :	othe 222. FION RMAT	.359 METI	id re	enti gion T50	ty 92 33.					
(i	ix)	(A) (B) (C)	LOC	E/KE ATIC	N: 3 ICAT	other 348 CION CMATI	393 METH	ide reg	entit gion T500	y 10 160.		,			
(i	х) 1	(A) (B) (C)	LOC.	ATIO NTIF	N: 1 ICAT	ther 89 ION MATI	229 METH	ide reg	ntit ion T500	y 97 14					
		(B) (C) (D)	NAMI LOCA I DEN	ATIO NTIF: ER II	N: 1: ICAT: NFOR	MATI	196 METHO ON:	OD: sco seq	re 4 QFI	Heij: .4 LLGT'	TSVV				
GACTGATT	rc G	AGTT	TCC	GG T	CAGG'	TTAG	G CC	GGGG	GGGT	GCG	GTCC	TGG	TCGG	AAGGAG	60
GTGGAGAG	rc G	GGGG	STCAC	CC A	GCC'	TATC	C TT	GGCG	CCAC	AGT	CGGC	CAC	CGGG	GCTCGC	120
CGCCGTC A	ATG Met	GAG Glu	AGC Ser	GGA Gly -20	GGG Gly	CGG Arg	CCC Pro	TCG Ser	CTG Leu -15	TGC Cys	CAG Gln	TTC Phe	ATC Ile	CTC Leu -10	169
CTG GGC A Leu Gly T	ACC	ACC Thr	TCT Ser -5	GTG Val	GTC Val	ACC Thr	GCC Ala	GCC Ala 1	CTG Leu	TAC Tyr	TCC Ser	GTG Val 5	TAC Tyr	CGG Arg	217
CAG AAG G Gln Lys A	SCC (Ala .	CGG Arg	GTC Val	TCC Ser	CAA Gln	GAG Glu 15	CTC Leu	AAG Lys	GGA Gly	GCT Ala	AAA Lys 20	AAA Lys	GTT Val	CAT His	265
TTG GGT G Leu Gly G 25	GAA (Glu)	GAT Asp	TTA Leu	AAG Lys	AGT Ser 30	ATT Ile	CTT Leu	TCA Ser	GAA Glu	GST Xaa 35	CCA Pro	GGA Gly	AAA Lys	TGC Cys	313
GTG CCT T Val Pro T 40	AT (GCT Ala	GTT Val	ATA Ile 45	GAA Glu	GGA Gly	GCT Ala	GTG Val	CGG Arg 50	TCT Ser	GTT Val	AAA Lys	GAA Glu	ACG Thr 55	361

391

CTT AAC AGC CAG TTT GTG GAA AAC TGC AAG Leu Asn Ser Gln Phe Val Glu Asn Cys Lys 60 65

(2) INFORMATION FOR SEQ ID NO: 194:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 459 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 269..342
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 2..75 id R33746 est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 391..459
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 94

region 124..192 id R33746

204

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 344..391
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95

region 78..125 id R33746

est

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 397..453
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.1

seq IYIICFXLPPLFS/FN

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 194:

ATATATAAAT GTTTCATGTT ATTGGTTTTG TACCTAGTCC TTTGCATGGA TATATAGGTA 60
CCTAATGAAA ATCGAGGATC AGTGTATGAC AAATCTCCCA TCCTCCCCTT TCCTTATTGC 120

CTGTGTCGGC AATAGGAAGT AGAATAGTTG TGTGTTGTTT ACTTACTTGT CTG	GTTTTAGA	180
GAGATTTCTA TTTTTGGTAG GGGAATATTC TAATATGTTT TCATATCTTT ATT	TTCATTTT :	240
GTAGTCTTTT GCATGGCTAT GTAGGGACCT AATGAAAGTC GAGTTTCATA ATA	ATGACAGC (300
TCACDTCTTT TCCTACATAT TTCCTCACTT AGCAGTAGCT WGNKAGTTAT KTT	IGTGGTTA 3	360
TTTTATTTCA TTCTCTAGGA TCTATTCCAT TTGNNG ATG CAA GTG TGT AG Met Gln Val Cys Ar -1	rg Cys	114
ATA TAT ATC ATT TGC TTC TWT CTT CCG CCA TTA TTT TCC TTT AA Ile Tyr Ile Ile Cys Phe Xaa Leu Pro Pro Leu Phe Ser Phe As -10	AC 4	159

(2) INFORMATION FOR SEQ ID NO: 195:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 193 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 44..193
 - (C) IDENTIFICATION METHOD: fasta
 - (D) OTHER INFORMATION: identity 96.1 region 1..152 id HSU78678 vrt

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 112..193
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 90..171

id N41898

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 112..193
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 95..176

id H69272

est

(ix) FEATURE:

11.0.131,00,00		161	
(B) (C)	NAME/KEY: other LOCATION: 112193 IDENTIFICATION METHO OTHER INFORMATION:	DD: blastn identity 97 region 39120 id N20619 est	
(B) (C)	NAME/KEY: sig_peptid LOCATION: 4488 IDENTIFICATION METHO OTHER INFORMATION:		
(xi) SEQU	ENCE DESCRIPTION: SEQ) ID NO: 195:	
GGGAGGGCTA GGCT	GTGCAT CCCTCCGCTC GCA	ATTGCAGG GAG ATG GCT CA Met Ala Gl -15	
CTT CTT CTG AGG Leu Leu Leu Arg -10	TTC CTG GCC TCT GTC Phe Leu Ala Ser Val -5	ATC TCC AGG AAG CCC TC Ile Ser Arg Lys Pro Se 1	T CAR 103 r Gln 5
GGT CAG TGG GCC Gly Gln Trp Ala	ACC CCT CAC TTC CAG Thr Pro His Phe Gln 10	AGC CCT GCA GAC CCC AC. Ser Pro Ala Asp Pro Th 15	r Met
CAG TCC TGG TGG Gln Ser Trp Trp 25	CCT GAC TGT AAC ACC Pro Asp Cys Asn Thr 30	CAA CCC AGC CCG GAC Gln Pro Ser Pro Asp 35	193
(2) INFORMATION	FOR SEQ ID NO: 196:		
	NCE CHARACTERISTICS: LENGTH: 280 base pai	rs	
	TYPE: NUCLEIC ACID STRANDEDNESS: DOUBLE		
	TOPOLOGY: LINEAR		
(ii) MOLE	CULE TYPE: CDNA		
	INAL SOURCE:		
	ORGANISM: Homo Sapie TISSUE TYPE: Cancero		

(ix) FEATURE: (A) NAME

(A) NAME/KEY: other
(B) LOCATION: 111..277

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99 region 3..169 id AA149704

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 143..262

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 3.9 seq FLWLITRPOPVLP/LL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 196:

AAGTCCTAGG AGCTGTGGAA AGAGTAGAAG TGCCTGAATG TGGTGCTGAA TCAATACAGC 60 CAGCTGTGAG GGGAGCACTT CCTGGACCCA GGAAGGGAGA GTCTTCTTCC AAGGTCTGAA 120 TTTCCTGCTG CTGTTCACAA AG ATG CTT TTT ATC TTT AAC TTT TTG TTT TCC Met Leu Phe Ile Phe Asn Phe Leu Phe Ser -35 CCA CTT CCG ACC CCG GCG TTG ATC TGC ATC CTG ACA TTT GGA GCT GCC Pro Leu Pro Thr Pro Ala Leu Ile Cys Ile Leu Thr Phe Gly Ala Ala 220 -30 -25 ATC TTC TTG TGG CTG ATC ACC AGA CCT CAA CCC GTC TTA CCT CTT 268 Ile Phe Leu Trp Leu Ile Thr Arg Pro Gln Pro Val Leu Pro Leu Leu -5 GAC CTG AAC CKG 280 Asp Leu Asn Xaa

(2) INFORMATION FOR SEQ ID NO: 197:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 443 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 323..443
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 2..122 id R84934

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 323..390
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 1..68 id AA020870 est

WO 99/06550 163

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 215 base pairs
(B) TYPE: NUCLEIC ACID
(C) STRANDEDNESS: DOUBLE
(D) TOPOLOGY: LINEAR

(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 373443 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 92 region 52122 id AA020870 est	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(407438) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 90 region 4273 id AA187611 est	
(ix)	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 297434 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.9 seq SHMLQLLPSKALC/LF	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 197:	
TTTGTGGGCT	CCTCTTTGGG GTGACCACTG CTTTCAAAGC CATCTGCCAA GGCTCTCCAG	60
GGCAGGACCT	GACTGGTGGG GAATGAGTGT TCAGAAGCCT TGGGAGAGGC CAAAGAGCCA 1	20
TTCTAGGATG	RTCKGAGGAA AACCTTCCTG CAGAGGCCAG AAACCTTGAG CTTAGGTGCC 1	80
TGGGGACCAG	CTTCGACATT CTCTCCAGTT TCTGATTCTA ATTTTTGCCA CGTGTCACAA 2	4 (
CTTTTCCAGT	CTCTGAGAAG GTCCCAGVCT TTCTCAAATA TTCTGATTTT GAAAAT ATG 2 Met	99
TAT CCA AAG Tyr Pro Ly -45	S TGG GAG GCC CCT GTG ACA TTT TGC CAA CTT AAA CGA GAA S Trp Glu Ala Pro Val Thr Phe Cys Gln Leu Lys Arg Glu -40 -35 -30	47
AAA GAC CC Lys Asp Pr	C CCG CAC CCG GCA CAC TCC CCC TTC CTC CAG CCC CGC TTC Pro His Pro Ala His Ser Pro Phe Leu Gln Pro Arg Phe -25 -20 -15	95
	G CTC CAG CTG CTG CCC AGT AAA GCC CTG TGC CTT TTT TTC Leu Gln Leu Leu Pro Ser Lys Ala Leu Cys Leu Phe Phe -10 -5 1	43
(2) INFORM	ATION FOR SEQ ID NO: 198:	

(ii) MOLECULE TYPE: CDNA

```
(vi) ORIGINAL SOURCE:
             (A) ORGANISM: Homo Sapiens
             (F) TISSUE TYPE: Cancerous prostate
       (ix) FEATURE:
             (A) NAME/KEY: other
             (B) LOCATION: 42..151
             (C) IDENTIFICATION METHOD: blastn
             (D) OTHER INFORMATION: identity 98
                                     region 1..110
                                     id AA121585
                                     est
       (ix) FEATURE:
             (A) NAME/KEY: other
             (B) LOCATION: 143..214
             (C) IDENTIFICATION METHOD: blastn
             (D) OTHER INFORMATION: identity 95
                                     region 101..172
                                     id AA121585
                                     est
       (ix) FEATURE:
             (A) NAME/KEY: other
             (B) LOCATION: 42..136
             (C) IDENTIFICATION METHOD: blastn
             (D) OTHER INFORMATION: identity 100
                                     region 1..95
                                     id AA100539
                                     est
       (ix) FEATURE:
            (A) NAME/KEY: other
            (B) LOCATION: 143..214
            (C) IDENTIFICATION METHOD: blastn
            (D) OTHER INFORMATION: identity 95
                                     region 100..171
                                     id AA100539
                                     est
      (ix) FEATURE:
            (A) NAME/KEY: sig_peptide
            (B) LOCATION: 36..167
            (C) IDENTIFICATION METHOD: Von Heijne matrix
            (D) OTHER INFORMATION: score 3.7
                                    seq LAERLGLFEELWA/AQ
      (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 198:
ACTGTTTGAG GATGTAGGCA CTGGTGTGAA GGAAC ATG GCC CTG TAT CAG AGG
                                                                     53
                                        Met Ala Leu Tyr Gln Arg
TGG CGG TGT CTC CGG CTC CAA GGT TTA CAG GCT TGC AGG CTA CAC ACG
Trp Arg Cys Leu Arg Leu Gln Gly Leu Gln Ala Cys Arg Leu His Thr
```

GCA GTT GTG TCG ACC CCT CCA CGC TGG TTG GCA GAG CGG CTT GGC CTT
Ala Val Val Ser Thr Pro Pro Arg Trp Leu Ala Glu Arg Leu Gly Leu
-20

TTT GAG GAG CTG TGG GCT CAG GTA AAG AGA TTA GCA AGC ATG GCA
Phe Glu Glu Leu Trp Ala Ala Gln Val Lys Arg Leu Ala Ser Met Ala
-5

CAG AAG GAA CCC CAG ACG

TOTAL CAG ACG

215

Gln Lys Glu Pro Gln Thr 15

(2) INFORMATION FOR SEQ ID NO: 199:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 280 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 57..276
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95

region 22..241

id C16912

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 172..260
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 91

region 64..152

id T68684

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 132..164
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96

region 26..58

id T68684

est

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 98..166
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 13.8

seq XGLLLFLLPGSLG/AE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 199:

AGAGAGAGA ACTGGGGTCT CCAGTCACGG GAGCCAGGAG CCGCCAGGAG CCGCAGSAGG AAGGGAGCGA GGCTGAAGGG AACGTCGTCC TCTCAGC ATG GGG GTC CCG CGG CCT Met Gly Val Pro Arg Pro CAG CCC TGG GCG STG GGG CTC CTG CTC TTT CTC CTT CCT GGG AGC CTG Gln Pro Trp Ala Xaa Gly Leu Leu Phe Leu Leu Pro Gly Ser Leu 163 -15 GGC GCA GAA AGC CAC CTC TCC CTC CTG TAC CAC CTT ACC GCG GTG TCC Gly Ala Glu Ser His Leu Ser Leu Leu Tyr His Leu Thr Ala Val Ser TCG CCT GCC CCG GGG ACT CCT GCC TTC TGG GTG TCC GGC TGG CTG GGC Ser Pro Ala Pro Gly Thr Pro Ala Phe Trp Val Ser Gly Trp Leu Gly 259 20 25 30 CCG CAG CAG TAC CCG AGC CAK 280 Pro Gln Gln Tyr Pro Ser Xaa 35

(2) INFORMATION FOR SEQ ID NO: 200:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 354 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 2..249
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 5..252 id C18087

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 166..350
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 20..204 id AA018305 est
- (ix) FEATURE:

WO 99/06550 PCT/IB98/01232

			(B) (C)	NAME LOCA I DEN OTHE	ATION ITIFI	: 18 :CATI	373 ON M	ETHO	iden regi	itity	/ 94 122	:05			
	()	ix) 1	(B) (C)	JRE: NAME LOCA IDEN OTHE	ATION ITIFI	: 18 :CATI	113 ON M	ETHO	iden regi	tity	, 96 332	02			
	(i)	ix) i	(B) (C)	JRE: NAME LOCA IDEN OTHE	TION TIFI	: 15 CATI	01 ON M	ETHO	iden regi	tity	100				
	(i	ix) I	(B) (C)	JRE: NAME LOCA IDEN OTHE	TION TIFI	: 15 CATI	83 ON M	ETHO	iden regi	tity	7 95 .21	92			
	-		(B) (C) (D)	JRE: NAME LOCA IDEN OTHE	TION TIFI CR IN	: 28 CATI	ON M	62 METHO ON:	D: V scor seq	e 13 LVLA	3.4 LXLV	SAAL			
AAGO	CGCA	GC 1	rccc;	AGCC	GA G1	rccgr		et Al					o Ly	GG ATG	54
				CAA Gln											102
				GCG Ala											150
				AGT Ser 1											198

WO 99/06550 PCT/IB98/01232

GTA Val	CTC Leu	AAC Asn 15	TCA Ser	CAT His	ATT Ile	TCT Ser	ACC Thr 20	CCA Pro	AAT Asn	GTG Val	AAT Asn	GCT Ala 25	TTA Leu	ACA Thr	CAT His	246
GAA Glu	AAC Asn 30	CAA Gln	ACC Thr	AAA Lys	CCT Pro	TCT Ser 35	ATT Ile	TCC Ser	CAA Gln	ATC Ile	AGC Ser 40	ACC Thr	ACC Thr	CTC Leu	CCT Pro	294
CCC Pro 45	AYR Xaa	NCG Xaa	AGT Ser	ACC Thr	AAG Lys 50	VNA Xaa	AGT Ser	GGA Gly	GGA Gly	GCA Ala 55	TYT Xaa	GTG Val	GTC Val	CCT Pro	CAT His 60	342
	TCG Ser															354

(2) INFORMATION FOR SEQ ID NO: 201:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 334 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 170..322
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 117..269 id HSC3DG011

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 53..184
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 91

region 1..132 id HSC3DG011

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(177..209)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 90

region 337..369

id H41589

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 137..223

169

(C) IDENTIFICATION METHOD: Von Heijne matrix
(D) OTHER INFORMATION: score 13
seq LLLVLLLVTRXRS/MP

...

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 201:

AATTTGTGCG GCGCTGGTCC CCTCAGAGGG TTCCTGCTGC TGCCGGTGCC TTGGACCCTC 60

CCCCTCGCTT CSNGTTCTAC TGCCCCAGGA GCCCGGCGGG TCCGGGACTC CCGKCCGTGC 120

CGGTGCGGCC GCCGCC ATG TGG CTG TGG GAG GAC CAG GGC GGC CTC CTG GGC 172

Met Trp Leu Trp Glu Asp Gln Gly Gly Leu Leu Gly

-25

-20

CCT TTC TCC TTC CTG CTG CTA GTG CTG CTG CTG GTG ACG CGG ASC CGG
Pro Phe Ser Phe Leu Leu Leu Val Leu Leu Val Thr Arg Xaa Arg
-15 -10 -5

TCA ATG CCT GCC TCC TCA CCG GCA GCC TCT TCG TTC TAC TGC GCG TCT

Ser Met Pro Ala Ser Ser Pro Ala Ala Ser Ser Phe Tyr Cys Ala Ser

1 10 15

TCA GCT BTG AGC CGG TGC CCT CTT GCA GGG CCC TGC AGG TGC TCA AGC
Ser Ala Xaa Ser Arg Cys Pro Leu Ala Gly Pro Cys Arg Cys Ser Ser
20 25 30

CCC GGG ACC GCA TTT CTG
Pro Gly Thr Ala Phe Leu
35

(2) INFORMATION FOR SEQ ID NO: 202:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 281 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 24..280

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 28..284 id R02745

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 3..176

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 6..179 id T84331 est

(ix) FEATURE:

- (A) NAME/KEY: other
 (B) LOCATION: 172..280
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 176..284 id T84331 est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 27..280
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100 region 1..254 id AA017512 est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 27..280
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 1..254 id N95074 est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 173..280
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 146..253 id N75564 est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 65..151
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 40..126 id N75564

est

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 27..66
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 92

region 1..40 id N75564

(ix) FEATURE:

(A) NAME/KEY: sig_peptide (B) LOCATION: 36..119

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 11.6 seq LLLLVQLLRFLRA/DG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 202:

ATTTCTTCCC CCCGAGCTGG GCGTGCGCGG CCGCA ATG AAC TGG GAG CTG CTG Met Asn Trp Glu Leu Leu -25 CTG TGG CTG CTG CTG TGC GCG CTG CTC CTG CTC TTG GTG CAG CTG 101 Leu Trp Leu Leu Val Leu Cys Ala Leu Leu Leu Leu Leu Val Gln Leu -20 CTG CGC TTC CTG AGG GCT GAC GGC GAC CTG ACG CTA CTA TGG GCC GAG 149 Leu Arg Phe Leu Arg Ala Asp Gly Asp Leu Thr Leu Leu Trp Ala Glu TGG CAG GGA CGC CCA GAA TGG GAG CTG ACT GAT ATG GTG GTG TGG 197 Trp Gln Gly Arg Arg Pro Glu Trp Glu Leu Thr Asp Met Val Val Trp 15 20 GTG ACT GGA GCC TCG AGT GGA ATT GGT GAG GAG CTG GCT TAC CAG TTG Val Thr Gly Ala Ser Ser Gly Ile Gly Glu Glu Leu Ala Tyr Gln Leu 35 TCT AAA CTA GGA GTT TCT CTT GTG CTG TCA GCC AGG 281 Ser Lys Leu Gly Val Ser Leu Val Leu Ser Ala Arg

(2) INFORMATION FOR SEQ ID NO: 203:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 344 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 163..344

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 35..216 id T86663

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 163..278

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

172

region 43..158 id AA055880 est

		FEATUR	

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 177..236
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 11.2

seq AFLLLVALSYTLA/RD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 203:

AGAAGATAAT CACTTGGGGA AAGGAAGGTT CGTTTCTGAG TTAGCAACAA GTAAATGCAG CACTAGTGGG TGGGATTGAG GTATGCCCTG GTGCATAAAT AGAGACTCAG CTGTGCTGGC ACACTCAGAA GCTTGGACCG CATCCTAGCC GCCGACTCAC ACAAGGCAGA GTTGCC ATG 179 GAA AAA ATT CCA GTG TCA GCA TTC TTG CTC CTT GTG GCC CTC TCC TAC 227 Glu Lys Ile Pro Val Ser Ala Phe Leu Leu Leu Val Ala Leu Ser Tyr -10 ACT CTG GCC AGA GAT ACC ACA GTC AAA CCT GGA GCC AAA AAG GAC ACA 275 Thr Leu Ala Arg Asp Thr Thr Val Lys Pro Gly Ala Lys Lys Asp Thr AAG GAC TCT CGA CCC AAA CTG CCC CAG ACC CTC TCC AGA GGT TGG GGT 323 Lys Asp Ser Arg Pro Lys Leu Pro Gln Thr Leu Ser Arg Gly Trp Gly 20 GAC CAA CTC ATC TGG ACA CGG 344

(2) INFORMATION FOR SEQ ID NO: 204:

Asp Gln Leu Ile Trp Thr Arg

30

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 312 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (iz) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 171..312
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95 region 33..174 id T86663

(ix)	FEAT	FEATURE:										
	(A)	NAME/KEY: other										
	(B)	LOCATION: 171288										
	(C)	IDENTIFICATION METHOD: blastn										
	(D)	OTHER INFORMATION: identity 97										

region 41..158 id AA055880 est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 127..246
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 11.2 seq AFLLLVALSYTLA/RD
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 204:

AAGAT'	rcaca	AGGC	CAAC	AG A	CAAC	CCAA	A GT	CATT	'AAGC	CAT	GAGA	GTG	GAAT	GAATCI	60
ATGAA	AACTC	AATG	AAGA	CA G	AACA	AGAG	a aa	AATC	TTTT	CAG	CCAC	GAT	GAAT:	raggro	120
AACAA		TCA . Ser .			Thr					Ser					168
GAG A' Glu II						Glu									216
CTC CT Leu Le -10															264
CCT GG Pro G			Lys												312

(2) INFORMATION FOR SEQ ID NO: 205:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 326 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other(B) LOCATION: 96..165
 - (C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 364..433

id AA100852 est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 45..95

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 314..364

id AA100852

est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 14..46

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 282..314

id AA100852

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 96..202

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 94

region 65..171

id AA113841

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 31..95

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 1..65

id AA113841

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 290..324

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 326..360

id AA133048

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 158..191

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 2..35 id AA133048

est

(ix) FEATURE:

(A) NAME/KEY: other

			(C)	LOCA IDEN OTHE	TIFI	CATI	ON N	иетно	ider regi	tity	97 12	22				
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				NAME				<u>,</u>								
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										on 3 A161	65	407				
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	t×	(i) S	EQUE	NCE	DESC	RIPT	:NOI	SEC	O ID	NO:	205:	;				
AC A	ATG C	CAG I	TT C	SNA A	ACG I	GG (GCT A	ACT T	rcc 1	CA 1	rcg (CAG C	cc c	ברה יו	יייכ	47
			he >	(aa 1				Chr S	Ser S			Gln E	Pro P	Ala I		3,
			-	-50				-	-45				-	-40		
TGG	TCT	TTG	CTC	TTG	GTT	TCC	TGG	GCT	GCT	ATG	GTG	CTA	AGA	CTG	AGA	95
Trp	Ser	Leu	_35	Leu	val	Ser	Trp	A1a -30	Ala	Met	Val	Leu	Arg -25	Leu	Arg	
		mcm.	000	0.00	cm.c											
												CTC Leu				143
	•	-20					-15					-10				
ATT	GCT	GAG	GTT	GCA	GCT	GCT	GTG	GTC	GCC	TTG.	GTG	TAC	ANC	ACA	ATG	191
Ile	Ala	Glu	Val	Ala	Ala	Ala	Val	Val	Ala	Leu	Val	Tyr	Xaa	Thr	Met	171
	- 5					1				5					10	
BCT	GAG	CAC	TTC	CTG	ACG	TTG	CTG	GTA	GTG	ССТ	GCC	ATC	AAG	AAA	GAT	239

Xaa Glu His Phe Leu Thr Leu Leu Val Val Pro Ala Ile Lys Lys Asp 15

TAT GGT TCC CAG GAA GAC TTC ACT CAA GTG TKG AAC ACC ACC ATG AAA 287 Tyr Gly Ser Gln Glu Asp Phe Thr Gln Val Xaa Asn Thr Thr Met Lys 30 35

GGG CTC AAG TGC TGT GGC TTC ACC AAC TAT ACG GAC TGG 326 Gly Leu Lys Cys Cys Gly Phe Thr Asn Tyr Thr Asp Trp 45 50

(2) INFORMATION FOR SEQ ID NO: 206:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 335 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 140..276
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 147..283

id N36076

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 32..140
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 40..148

id N36076

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 287..333
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 296..342

id N36076

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 1..33
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 90 region 8..40 id N36076

177

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 2..333
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 9..340 id N95074

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 2..333
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 9..340

id AA017512

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 140..333
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 146..339

id W04626

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 5..140
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 12..147

id W04626

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 45..334
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 47..336

id H27747

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 1..34
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 1..34

id H27747

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 3..86
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.5

seq LLLLVHLLRFLRA/DG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 206:

AA	ATG Met	AAC Asn	TGG Trp	GAG Glu -25	CTG Leu	CTG Leu	CTG Leu	Trp	CTG Leu -20	CTG Leu	GTG Val	CTG Leu	Cys	GCG Ala -15	CTG Leu	` 47
CTC Leu	CTG Leu	CTC Leu	Leu -10	GTG Val	CAT His	CTG Leu	CTG Leu	CGC Arg -5	TTC Phe	CTG Leu	AGG Arg	GCT Ala	GAC Asp	GGC Gly	GAC Asp	95
CTG Leu	ACG Thr 5	CTA Leu	CTA Leu	TGG Trp	GCC Ala	GAG Glu 10	Trp	CAG Gln	GGA Gly	CGA Arg	CGC Arg 15	CCA Pro	GAA Glu	TGG Trp	GAG Glu	143
CTG Leu 20	ACT Thr	GAT Asp	ATG Met	GTG Val	GTG Val 25	TGG Trp	GTG Val	ACT Thr	GGA Gly	GCC Ala 30	TCG Ser	AGT Ser	GGA Gly	ATT Ile	GGT Gly 35	191
GAG Glu	GAG Glu	CTG Leu	GCT Ala	TAC Tyr 40	CAG Gln	TTG Leu	TCT Ser	AAA Lys	CTA Leu 45	Gly	KTT Xaa	TCT Ser	CTT Leu	GTG Val 50	CTG Leu	239
TCA Ser	GCC Ala	AGA Arg	AGA Arg 55	GTG Val	CAT His	GAG Glu	CTG Leu	GAA Glu 60	AGG Arg	GTG Val	AAA Lys	AGA Arg	AGA Arg 65	TGC Cys	CTA Leu	287
GAG Glu	AAT Asn	GGC Gly 70	AAT Asn	TTA Leu	ARA Xaa	GAA Glu	AAA Lys 75	GAT Asp	ATA Ile	CTT Leu	GTT Val	TTG Leu 80	CCC Pro	CTT Leu	GGG Gly	335

(2) INFORMATION FOR SEQ ID NO: 207:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 347 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (3) LOCATION: 53..162
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 424..533 id N80896 est
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (3) LOCATION: complement(283..318)

				OTHE					ider regi	ntity	7 100 342					
	(i	ix) I	(B) (C)	JRE: NAME LOCA I DEN OTHE	TION	1: 29 CATI	33 ON N	ETHO	ider regi	tity	100					
			(B) (C) (D)	NAME LOCA IDEN OTHE	TION TIFI CR IN	: 12 CATI	202 ION N MATIC	272 METHO ON:	D: V scor seq	e 10 VSCI	.3 TLWS	SPGCV				
	()	(i) S	SEQUE	ENCE	DESC	CRIPT	rion:	: SE() ID	NO:	207:	:				
TGC	CTAT	rgc i	rtgto	GTGT <i>A</i>	AT G	rgtg:	rgcc:	г ст	STCT	rgct	CTC	TATO	CTC (CCAGO	CAGTGA	60
GAC	ATTGO	GAC (GTGTT	rtgci	C A	rgaa(GATG	C AG	CATAT	rggc	TTG	rctg:	rga (GCCCA	AGTGA	119
			TTT Phe													167
			GGG Gly													215
			CCT Pro													263
		Pro	CAG Gln l	Pro		Gln		Lys								311
			AGA Arg													347
(2)			(3)		CHARA STH:	ACTEI 461 JCLE:	RIST: base IC A	ICS: e pai								
			(\mathfrak{D})	TOPO	OLOG!	7 · T.	INFAI	₹ .								

(ii) MOLECULE TYPE: CDNA

WO 99/06550 PCT/IB98/01232

(vi)	(A) OR	L SOURCE GANISM: I SSUE TYPE	Homo Sap	oiens erous	pros	tate							
(ix)	(B) LO	ME/KEY: c CATION: 1 ENTIFICAT	168461 TION MET	HOD: ide reg id	HOD: blastn identity 99 region 47340 id N39924 est								
(ix)	(B) LOC (C) IDE	: ME/KEY: c CATION: 1 ENTIFICAT HER INFOR	.69370 'ION MET	HOD: ide reg	blas ntit ion R616	y 96 178.	. 379						
(ix)	(B) LOC (C) IDE	ME/KEY: o MATION: 3 MTIFICAT MER INFOR	59431 ION MET	ide: reg	blas ntit ion (y 100 369.							
(ix) E	(B) LOC	E/KEY: s ATION: 7 NTIFICAT ER INFOR	5158 ION MET	HOD: '	Von H re 9. LVXI	. 5							
(xi) S	SEQUENCE	DESCRIP	TION: S	EQ ID	NO:	208	:						
ACCATAGCAA #	ATTAAATO	AC TGCCA	TAAAG T	ATATT	TTAC	TCA	CAGG	ACA (GATT	ACAATA	60		
GCCTTGATAG A	AATC ATG Met	GCA TCC Ala Ser	AAA GG Lys G1 -25	G ATG y Met	CGC Arg	CAT His	TTT Phe -20	TGC Cys	TTG Leu	ATT Ile	110		
TCA GAG CAG Ser Glu Gln -15	TTG GTG Leu Val	TYC TTT Xaa Phe	Ser Le	T CTT u Leu	GCA Ala	ACA Thr -5	GCG Ala	ATT Ile	TTG Leu	GGA Gly	158		
GCA GTT TCC Ala Val Ser 1	TGG CAG Trp Gln	Pro Thr	AAT GG Asn Gl	A ATT y Ile 10	TTC Phe	TTG Leu	AGC Ser	ATG Met	TTT Phe 15	CTA Leu	206		
ATC GTT TTG Tie Val Leu	CCA TTG Pro Leu 20	GAA TCC Glu Ser	ATG GC Met Al	a His	GGG Gly	CTC Leu	TTC Phe	CAT His 30	GAA Glu	TTG Leu	254		
GGT AAC TGT Gly Asn Cys	TTA GGA	GGA ACA	TCT GT Ser Va	r GGA l Glv	TAT Tvr	GCT Ala	ATT	GTG Val	ATT	CCC	302		

35 40 45

ACC AAC TTC TGC AGT CCT GAT GGT CAG CCA ACA CTG CTT CCC CCA GAA 350 Thr Asn Phe Cys Ser Pro Asp Gly Gln Pro Thr Leu Leu Pro Pro Glu 50 55 CAT GTA CAG GAG TTA AAT TTG AGG TCT ACT GGC ATG CTC AAT GCT ATC 398 His Val Gln Glu Leu Asn Leu Arg Ser Thr Gly Met Leu Asn Ala Ile 70 CAA AGA TTT TTT GCA TAT CAT ATG ATT GAG ACC TAT GGA TGT GAC TAT 446 Gln Arg Phe Phe Ala Tyr His Met Ile Glu Thr Tyr Gly Cys Asp Tyr 90 TCC ACA AGT GGA CTG 461 Ser Thr Ser Gly Leu

(2) INFORMATION FOR SEQ ID NO: 209:

100

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 296 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement (31..239)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 93

region 3..211 id N27605 est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(2..111)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 1..110 id N78549

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 78..140
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 9.3 seq VLPVILLLLGAHP/SP
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 209:

AAG	AGCA	GAG	CCGG	AAGA	AG G	CGGG	ACGA	A CC	GGAA	GAGG	GTO	SAAAT	GCT	TTCG	GTAGGC	60
ACT	CCAC	GGC	TGTG.	AAG	ATG Met	GCG Ala -20	GCG Ala	GCT Ala	GCG Ala	TGG Trp	CTT Leu -15	CAG Gln	GTG Val	TTG Leu	CCT Pro	110
GTC Val -10	TTE	CTT Leu	CTG Leu	CTT Leu	CTG Leu -5	GGA Gly	GCT Ala	CAC His	CCG Pro	TCA Ser	CCA Pro	CTG Leu	TCG Ser	TTT Phe 5	TTC Phe	158
AGT Ser	GCG Ala	GGA Gly	CCG Pro 10	GCA Ala	ACC Thr	GTA Val	GCT Ala	GCT Ala 15	GCC Ala	GAC Asp	CGG Arg	TCC Ser	AAA Lys 20	TGG Trp	CAC His	206
RKT Xaa	CCG Pro	ATA Ile 25	CCG Pro	TCG Ser	GGG Gly	AAA Lys	AAT Asn 30	Tyr	TTT Phe	AGT Ser	TTT Phe	GGA Gly 35	AAG Lys	ATC Ile	CTC Leu	254
TTC Phe	AGA Arg 40	AAT Asn	ACC Thr	ACT Thr	ATC Ile	TTC Phe	CTG Leu	AAG Lys	TTT Phe	GAT Asp	GGA Gly	GAA Glu	CGA Arg			296

(2) INFORMATION FOR SEQ ID NO: 210:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 468 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 118..281
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 78..241 id R57572

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 38..91
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 1..54 id R57572

- (ix) FEATURE:
 - (A) NAME/KEY: other (B) LOCATION: 90..122
 - (C) IDENTIFICATION METHOD: blastn

WO 99/06550 PCT/IB98/01232

(D) OTHER INFORMATION: identity 93 region 52..84 id R57572 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 117..272 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 94 region 59..214 id W55468 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 273..328 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 91 region 214..269 id W55468 est (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 130..456 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 9.1 seq LVLAVLFFHQLVG/DP (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 210: ACTITGICAT TCAGCTGCCT GCTGCCTCCG CAGCGTCCCC CCAGCTCTCC CTGTGCTAAC TGCCTGCACC TTGGACAGAG CGGGTGCGCA AATCAGAAGG ATTAGTTGGG ACCTGCCCTT GGCGACCCC ATG GCA TCC CCC AGA ACC GTA ACT ATT GTG GCC CTC TCA GTG Met Ala Ser Pro Arg Thr Val Thr Ile Val Ala Leu Ser Val -105 GCC CTG GGA CTC TTC TTT GTT TTC ATG GGG ACT ATC AAG CTG ACC CCC 219 Ala Leu Gly Leu Phe Phe Val Phe Met Gly Thr Ile Lys Leu Thr Pro -90 -85 AGG CTC AGC AAG GAT GCC TAC AGT GAG ATG AAA CGT GCN NAC AAG AGC 267 Arg Leu Ser Lys Asp Ala Tyr Ser Glu Met Lys Arg Ala Xaa Lys Ser -70 TAT GTT CGA GCC CTC CCT CTG CTG AAG AAA ATG GGG ATC AAT TCC ATT 315 Tyr Val Arg Ala Leu Pro Leu Leu Lys Lys Met Gly Ile Asn Ser Ile CTC CTC CGA AAA AGC ATT GGT GCC CTT GAA GTG GCC TGT GGC ATC GTC 363 Leu Leu Arg Lys Ser Ile Gly Ala Leu Glu Val Ala Cys Gly Ile Val -40 ATG ACC CTT GTG CCT GGG CGT CCC AAA GAT GTG GCC AAC TTC TTC CTA Met Thr Leu Val Pro Gly Arg Pro Lys Asp Val Ala Asn Phe Phe Leu

(2) INFORMATION FOR SEQ ID NO: 211:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 225 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 88..221
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 84..217 id AA021055

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 5..74
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 1..70 id AA021055 est
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 88..221
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 84..217 id W98068

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 5..74
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 1..70 id W98068

- (ix) FEATURE:
 - (A) NAME/KEY: other

(B) LOCATION: 88191 (C) IDENTIFICATION METH (D) OTHER INFORMATION:	OD: blastn identity 99 region 11114 id AA059040 est	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptic (B) LOCATION: 91204 (C) IDENTIFICATION METHO (D) OTHER INFORMATION:</pre>	OD: Von Heijne matrix	
(xi) SEQUENCE DESCRIPTION: SEC	Q ID NO: 211:	
CATAAAATTT GAGGATATCA GCTGATTATT TT	TTCTTCCM ASAATGAAAA TCAAGCAGAA	60
TTGATTCCTA CACGAAAAAA AAGCACACGA ATG	G CCA AAC CTT TCC TTT GGT GGA 1: t Pro Asn Leu Ser Phe Gly Gly -35	14
CTG GAC ACT AAC CAG ATG AGA GTA AAT Leu Asp Thr Asn Gln Met Arg Val Asn -30 -25	TTC TTA TCC GTG GAC GTA TGT Phe Leu Ser Val Asp Val Cys -20 -15	62
AAG CTA CTG CTG CTG TGT GCT CTC CAC Lys Leu Leu Leu Cys Ala Leu His -10	AGC CAT ATT TAT TGT ATT AAA Ser His Ile Tyr Cys Ile Lys -5 1	10
CAA TCA GCA CTT CGG Gln Ser Ala Leu Arg 5	22	25
(2) INFORMATION FOR SEQ ID NO: 212:		
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 470 base pai (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR		
(ii) MOLECULE TYPE: CDNA		
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapie (F) TISSUE TYPE: Normal</pre>		
	DD: blastn identity 97 region 115359 id R67703 est	

(ix) FEATURE:

```
(A) NAME/KEY: other
```

(B) LOCATION: 23..135

(C) IDENTIFICATION METHOD: blastn
(D) OTHER INFORMATION: identity 92

region 5..117 id R67703

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 134..318
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 115..299

id H42383

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 20..135
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 2..117

id H42383

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 193..383
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 87..277

id W90193

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 134..192
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 29..87

id W90193

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 417..454
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 92

region 314..351

id W90193

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 288..470
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 1..183

id R53752

187

(iх)	FEA'	TURE:	
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(A) NAME/KEY: sig_peptide

(B) LOCATION: 258..422

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 8.8

seq XXLLLLNVGQLLA/QT

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 212:

AACCCAC	GGT GGG	GGGAGC	G CGGCC1	ATGGC G	CTCCT	GCTT	TCGGT	rgctgo	GTGT	ACTGCT	60
GGGCGGC1	TTC TTC	CGCGCTC	G TGGGG	TTGGC C	AAGCT	CTCG	GAGG	AGATC1	CGGC	rccagt	120
TTCGGAG	CGG RTG	RAATGC	C CTGTT	CGTGC A	GTTTG	CTGA	TGTGT	гтсссо	CTGA	AGGTAT	180
TTGGCTAG	CCA GCC	CAGATCC	C CTGAAG	CTACC A	AATAG	CTGT	GGGC1	TTTCTC	GAAC:	rgctgg	240
CTGGGTT(GCT GCT	M	TG GGC (et Gly I 55		Met 1				Ser A		290
TTG TTC Leu Phe											338
CTG AAA Leu Lys		r Leu			e Pro				s Leu		386
TDN CTG Xaa Leu											434
GTG GTC Val Val	Arg Pr	o Thr	Arg Lys		r Leu						470

(2) INFORMATION FOR SEQ ID NO: 213:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 354 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Cancerous prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 4..55
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98 region 19..70

id T18977 est

```
(ix) FEATURE:
```

- (A) NAME/KEY: other
- (B) LOCATION: 141..195
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 90

region 157..211

id T18977

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 92..137
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 93

region 109..154

id T18977

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 245..355
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 1..111

id HSC12A111

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 321..355
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 1..35

id W73324

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 133..345
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 8.6

seq VVXFLLLLAXLIA/TY

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 213:

AAGCAGCTTC CAGGATCCTG AGATCCGGAG CAGCCGGGGT CGGAGCGGCT CCTCAAGAGT

TACTGATCTA TNNATGGCAG AGAAAAAAA ATTGTGACCA GAGACGTGTA GCAATGAACA 120

AGGAACRTCA TA ATG RWN NNK TTC ACA GAC CCC TCT TCA GTG AAT GAA AAG 171 Met Xaa Xaa Phe Thr Asp Pro Ser Ser Val Asn Glu Lys

AAG AGG AGG GAG CAG GAA AGG CAG AAT ATT GTC CTG TGG AGA CAG 219 Lys Arg Arg Glu Arg Glu Arg Gln Asn Ile Val Leu Trp Arg Gln -55 -50

WO 99/06550 PCT/IB98/01232

									- '							
CCG Pro	CTC Leu	ATT Ile -40	ACC Thr	TTG Leu	CAG Gln	TAT Tyr	TTT Phe -35	TCT Ser	CTG Leu	GAA Glu	ATC Ile	CTT Leu -30	GTA Val	ATC Ile	TTG Leu	267
AAG Lys	GAA Glu -25	TGG Trp	ACC Thr	TCA Ser	AAA Lys	TTA Leu -20	TGG Trp	CAT His	CGT Arg	CAM Xaa	MGC Xaa -15	ATT Ile	GTG Val	GTG Val	WCT Xaa	315
				CTT Leu												354
(2)	(i)	.) SE	CQUEN (A) (B) (C) (D) MOLEC DRIGI	FOR NCE CLENG TYPE STRA TOPC CULE INAL ORGA TISS	CHARACTH: C: NU LNDED LOGY TYPE SOUR	ACTER 311 DOLEI DNESS : LI C: CI RCE:	RISTI base C AC : DC NEAF DNA	ICS: pai CID DUBLE	ens	state						
	(i	.x) E	(B) (C)	JRE: NAME LOCA IDEN OTHE	TION TIFI	: 18 CATI	93 ON M	1ETHC	ider regi		97 51	97				
	(i	.x) E	(B) (C)	JRE: NAME LOCA IDEN OTHE	TION TIFI	I: 24 CATI	92 ON M	293 METHO ON:	D: V	on H e 8. LLRG	6					
	()	(i) S	EQUE	ENCE	DESC	RIPT	CION:	: SE() ID	NO:	214:					
ACCI	TTC	rgg (STTG	AGCAT	rg go	CTGA	AGTG!	A CTO	CAGC	CCAT	GGG	AGGTT	TTC C	TAGO	GAGNAA	60
CAGO	CTC	CAC 7	TTGC:	rgcc	гс то	CTGC	TGA	A CTO	CCGT	GTGC	CGG	CAACO	CTG (CGA	CCAGAC	120
TCCI	GCC	rtc (GGAG	GGC	rg go	GCT	CCAG	G AC	CTGA	STGC	ccc	CCRNI	GT T	rgga <i>i</i>	AGGCGG	180
TGTO	ATA:	rgt (GCAC	AGAA	GC C	AAA?	AGCA	r TG	CTGG'	TATT	TCG	AAGG#	ACT (CTATO	CCAACC	240
YHTI	ATA'	r ATO	t Pro	G CT(C CT	A CGI	A GG g Gl g -10	y Le	G CTO	G TGO	S STO	C CAC a Glr	n Vai	CTO Le	G TGT	290
				CAT His												311

1

(2) INFORMATION FOR SEQ ID NO: 215:

```
(i) SEQUENCE CHARACTERISTICS:
```

- (A) LENGTH: 353 base pairs
- (B) TYPE: NUCLEIC ACID

5

- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Cancerous prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 121..355
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99 region 31..265

id T78247

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 121..355
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 6..240

id W17118

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 121..355
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 11..245

id N88433

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 121..336
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 32..247

id R35014

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 121..329
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99 region 9..217

id AA074562

(ix) FEA	TURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 159..218

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 8.4

seq AVVGCLLVPPAEA/NK

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 215:

AAGAGGCGGA GATGGCGGAG GGCGGTGGGA CGTGATGCGC GGGTCAGAGC CGGGCCTTGA 60 GAAGGAACTG GAGGCCCCTG GCAGCGGTGT CCCCTCGAGG ACCCCTCTGC CGGGCTCACC 120 AGGTGTCCGG CTTTGCTGGC CCAGCAAGCC TGATAAGC ATG AAG CTC TTA TCT TTG 176 Met Lys Leu Leu Ser Leu GTG GCT GTG GTC GGG TGT TTG CTG GTG CCC CCA GCT GAA GCC AAC AAG 224 Val Ala Val Val Gly Cys Leu Leu Val Pro Pro Ala Glu Ala Asn Lys -10 -5 AGT TCT GAA GAT ATC CRG TGC AAA TGC ATC TGT CCA CCT TAT AGA AAC Ser Ser Glu Asp Ile Xaa Cys Lys Cys Ile Cys Pro Pro Tyr Arg Asn ATC AGT GGG CAC ATT TAC AAC CAG AAT GTA TCC CAG AAG GAC TGC AAC 320 Ile Ser Gly His Ile Tyr Asn Gln Asn Val Ser Gln Lys Asp Cys Asn TGC CTG CAC GTG GTG GAG CCC ATG CCA GTG CCG 353 Cys Leu His Val Val Glu Pro Met Pro Val Pro 35 40

(2) INFORMATION FOR SEQ ID NO: 216:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 320 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 2..319
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 15..332 id HUMO85F04B

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 139..249

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100 region 187..297

id H85714

est

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 249..319

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 298..368

id H85714

est

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 86..148

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 92

region 133..195

id H85714

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 135..319

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 80..264

id R77008

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 86..319

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 94..327

id H49758

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 135..319

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 53..237

id AA056366

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 114..185

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 7.9

seq LLLPRVLLTMASG/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 216:

AATTGGCTGG CTCT	GGAGGC GCAGGTGG	TC CTTCTTCTAC	TGTCACATGG TGCGCGCTGT	60
TTTCTAATCA CGKG	GCTGCC ACCCAGGCC	CT CTCTGCTCCT	GTCKTKTGTT TGG ATG Met	116
			TTG CTA CCC CGA GTC Leu Leu Pro Arg Val -10	164
			CAG CCC TCG CCG GCC Gln Pro Ser Pro Ala 5	212
			GTC TCT GCA GCC TTT Val Ser Ala Ala Phe 25	260
			ATC GCC AGG GCC GTK Ile Ala Arg Ala Val 40	308
GGG GAG AAG CGG Gly Glu Lys Arg 45				320

(2) INFORMATION FOR SEQ ID NO: 217:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 384 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 121..381
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96

region 73..333 id H95186

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 72..133
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95 region 25..86 id H95186

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(1	\mathbf{x}) t	ĿΑ	Тι	JRE	:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 28..351
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 7.9

seq LLGLLSAEQLAEA/SV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 217:

ACGGGTGCCG GGTGGAGCGA ASACGGA ATG TGT CTC CTG CTG GGG GCC ACG GGC

Met Cys Leu Leu Gly Ala Thr Gly

-105

GTC GGG AAG ACG 'CTG CTG GTG AAA CGG CTG CAG GAG GTG AGC TCC CGG

Val Gly Lys Thr Leu Leu Val Lys Arg Leu Gln Glu Val Ser Ser Arg

-95

GAT GGG AAA GGC GAC CTG GGG GAG CCG CCC CCG ACA CGG CCC ACG GTG

ASD Gly Lys Cly Asp Leu Glo Gly Cac CCC CCC ACG GTG

150

Asp Gly Lys Gly Asp Leu Gly Glu Pro Pro Pro Thr Arg Pro Thr Val
-80 -75 -70

GGC ACC AAT CTT ACT GAC ATC GTG GCA CAG AGA AAG ATC ACC ATC CGG Gly Thr Asn Leu Thr Asp Ile Val Ala Gln Arg Lys Ile Thr Ile Arg -65 -55

GAG CTT GGG GGG TGC ATG GGC CCC ATC TGG TCC AGT TAC TAT GGA AAC
Glu Leu Gly Gly Cys Met Gly Pro Ile Trp Ser Ser Tyr Tyr Gly Asn
-50 -45 -40

TGC CGT TCT CTC CTG TTT GTG ATG GAC GCC TCT GAC CCC ACC CAG CTC

Cys Arg Ser Leu Leu Phe Val Met Asp Ala Ser Asp Pro Thr Gln Leu

-35 -20

TCT GCA TTM SGT GTG CAG CTC TTA GGT CTC CTT TCT GCA GAA CAA CTT
Ser Ala Xaa Xaa Val Gln Leu Leu Gly Leu Leu Ser Ala Glu Gln Leu
-15

GCA GAA GCA TCG GTG CTG ATA CTC TTC AAT AAA ATC GAC AAC
Ala Glu Ala Ser Val Leu Ile Leu Phe Asn Lys Ile Asp Asn

(2) INFORMATION FOR SEQ ID NO: 218:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 236 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other

(B) LOCATION: 94..197

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 92..195 id T93931 est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 2..45

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 1..44 id T93931 est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 53..97

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93

region 52..96 id T93931 est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 190..234

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 91

region 187..231 id T93931

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 138..196

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 241..299

id N25481

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 190..234

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93

region 292..336

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id N25481

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 94..211

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 65..182 id W19370

est.

WO 99/06550 196 (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 94..196 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 56..158 id N35539 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 190..234 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 93 region 151..195 id N35539 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 56..97 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 92 region 19..60 id N35539 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 94..193 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 96..195 id W87436 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 2..49 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 93 region 7..54 id W87436 est (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 75...197 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 7.7 seq LLCLGQLHHPGLG/RV (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 218:

AAAGTTTGTT CCCCGAGTTC GGAGCCTAGG AGCCCCCGC GGCTGCGGCG CAGGTGCCCT 60 CGGCCTTAGT CGGG ATG GAG CTG CCT GCK GTG AAC CTT GAA AGT GAT TCT 110 Met Glu Leu Pro Ala Val Asn Leu Glu Ser Asp Ser -40-35 -30

WO 99/06550 PCT/IB98/01232

CCT AGG TCA CTG GCT GCT GAC AAC CTG GGG CTG CAT TGT ATT CTC AGG
Pro Arg Ser Leu Ala Ala Asp Asn Leu Gly Leu His Cys Ile Leu Arg
-25

CTC CTA TGC CTG GGC CAA CTT CAC CAT CCT GGC CTT GGG CGT GTG GGC
Leu Leu Cys Leu Gly Gln Leu His His Pro Gly Leu Gly Arg Val Gly
-10

158

206

TGT GGC TCA GCG GGA CTC CAT CGA CGC CGG
Cys Gly Ser Ala Gly Leu His Arg Arg Arg
5
10

(2) INFORMATION FOR SEQ ID NO: 219:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 329 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 145..240
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95 region 99..194

id N28787

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 45..139
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 1..95 id N28787

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 253..326
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 207..280

id N28787

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 145..239
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 93

region 114..208

id AA102327 est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 59..139

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 30..110 id AA102327

est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 31..63

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 1..33 id AA102327

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 277..311

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 91

region 250..284 id AA102327

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 145..240

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 101..196 id AA019783

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 253..326

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 209..282 id AA019783

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 79..139

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 37..97 id AA019783

est

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 145..240

(C) IDENTIFICATION METHOD: blastn

. (D) OTHER INFORMATION: identity 95

region 115..210 id AA059290

est

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 41..139

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 90

region 13..111 id AA059290

est

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 253..319

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 92

region 223..289 id AA059290

est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 145..240

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 102..197

id H86516

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 253..326

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 210..283

id H86516

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 75..139

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 34..98 id H86516

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 171..323

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 7.6

seq PALILLFALGSLG/SG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 219:

200

GGT	GCTG'	rtg (CCAT	CATG	GC TO	GACCO	CCGA	c ccc	CCGG	FACC	CTC	GCTC	CTC (GATC	GAGGAC	120
GAC'	TTCA	ACT I	ATGG	CAGC	AA GO	CGTKC	GCY1	r cso	GCCA	CCGT	GCA	CATC		Met A		176
TTT Phe	CTG Leu	AGA Arg	AAA Lys	GTC Val -45	TAC Tyr	AGC Ser	ATT Ile	CTT Leu	TCT Ser -40	CTG Leu	CAG Gln	GTT Val	CTC Leu	TTA Leu -35	ACT Thr	224
ACA Thr	GTG Val	ACT Thr	TCA Ser -30	ACA Thr	GTT Val	TTT Phe	TTA Leu	TAC Tyr -25	TTT Phe	GAG Glu	TCT Ser	GTA Val	CGG Arg -20	ACA Thr	TTT Phe	272
GTA Val	CAT His	GAG Glu -15	AGT Ser	CCT Pro	GCC Ala	TTA Leu	ATT Ile -10	TTG Leu	CTG Leu	TTT Phe	GCC Ala	CTC Leu -5	GGA Gly	TCT Ser	CTG Leu	320
	TCG Ser 1															329

(2) INFORMATION FOR SEQ ID NO: 220:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 207 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUÉ TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 23..202
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 1..180 id W88492 est

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 25..111
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.6

seq PTLAIALAANAWA/FV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 220:

ACCATATGGG TGGTGTGGAT CGTC ATG TAT ACT TAC GGC AAC AAG CAG CAC 51 Met Tyr Thr Tyr Gly Asn Lys Gln His -25

									20	, 1						
	AGT Ser															99
	GCC Ala															147
	ACC Thr															195
	CGG Arg 30															207
(2)	INFO	ORMAT	rion	FOR	SEQ	ID I	10: 2	221:								
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 195 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(136167) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 90																
									•	ion 2 16276	239. 66	.270				
	(:	ix) l	(B) (C)	NAMI LOCA IDEI	ATION NTIF	Y: s: N: 70 ICAT: NFOR	016	65 METH	OD: 1	re 7						
	(xi)	SEQU	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	221	:				
ACT	TTCA	GTT	тсст	тстт	CC A	GCAC	GGAG	T AC	ACTG	СТСТ	GCC	TCCA	CTT	AGAT	TACTT	C 60

AGAAATGAA ATG CAG CAA ATA TTT ATC CAG CAG TGC AGG GAG TTG AAC TTT 111 Met Gln Gln Ile Phe Ile Gln Gln Cys Arg Glu Leu Asn Phe

TGG AGT CGG GAA CCT TGG ATT CTT GTT CTG GCT CTG CCA CTT ACT GTG

Trp Ser Arg Glu Pro Trp Ile Leu Val Leu Ala Leu Pro Leu Thr Val

-25

159

-30

-15

WO 99/06550	202	PCT/IB98/01232
	202	

Trp Pro Trp Leu Ser Pro Glu Ala Gln Pro Pro Leu 1 5 10	195
(2) INFORMATION FOR SEQ ID NO: 222:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 373 base pairs	
(B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:	
(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Cancerous prostate	
(ix) FEATURE:	
(A) NAME/KEY: other	
(B) LOCATION: 308370	
(C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98	
region 404466	
id AA158879	
est	
(ix) FEATURE:	
(A) NAME/KEY: sig_peptide	
(B) LOCATION: 110154	
(C) IDENTIFICATION METHOD: Von Heijne matrix	
(D) OTHER INFORMATION: score 7.5 seq AVLLALLMAGLAL/QP	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 222:	
NACTOCOTOC ACCANAGOO COCOMOTO A COMOTO	
AACTGGCTCC AGGAAACCCG CTGGTGTTGA CTGTGGGCAG TCCAGCCTCT CCCCATTTGA	
GGCCATATAA ANNACCTGAG GCCCTCTCCA CCACAGCCCA CCAGTGACC ATG AAG GCT	
Met Lys Ala -15	a
GTG CTG CTT GCC CTG TTG ATG GCA GGC TTG GCC CTG CAG CCA GGC ACT	166
Val Leu Leu Ala Leu Leu Met Ala Gly Leu Ala Leu Gln Pro Gly Thr	100
-10 -5 1	
GCC CTG CTG TGC TAC TCC TGG ARR GCC CAG GTG RGC AAC GAG GAC TGC	214
Ala Leu Leu Cys Tyr Ser Trp Xaa Ala Gln Val Xaa Asn Glu Asp Cys	
5 10 15 20	
CTG CAG GTG GAG AAC TGC ACC CAG CTG GGG GAG CAG TGC TGG ACC GCG	262
Leu Gln Val Glu Asn Cys Thr Gln Leu Gly Glu Gln Cys Trp Thr Ala	202
25 30 35	
CCC ATC CCC CCA CTT CCC CTC CTC CTC CTC	
CGC ATC CGC GCA GTT GGC CTC CTG ACC GTC ATC AGC AAA GGC TGC AGC Arg Ile Arg Ala Val Gly Leu Leu Thr Val Ile Ser Lys Gly Cys Ser	310

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										_				
				GAT Asp										358
	ACG Thr 70													373
(2)	INFO	ORMAT	CION	FOR	SEQ	ID N	10: 2	223:						
	(i	.) SE	(A) (B) (C)	ICE C LENG TYPE STRA TOPO	TH: : NU .NDED	249 CLEI NESS	base C AC : DC	pai ID UBLE						
	(i	.i) N	OLEC	ULE	TYPE	: CE	AAG							
	(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate													
	<pre>(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 1247 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100</pre>													
	<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 451 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 7.1 seq QACLLGLFALILS/GK</pre>													
	٠(>	(i) S	EQUE	ENCE	DESC	RIPT	CION:	SEÇ	DI	NO:	223:			
AGA				CAA Gln										48
				AGT Ser										96
				GTG Val 20										144
				GCC Ala										192

CTG GTG CAG GCT GTG TCG GAT CCC AGC TCT CCT CAA TAC GGA AAA TAC 240

204

Leu Val Gln Ala Val Ser Asp Pro Ser Ser Pro Gln Tyr Gly Lys Tyr 50 55 60

CTG ACC CGT Leu Thr Arg 65

249

- (2) INFORMATION FOR SEQ ID NO: 224:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 382 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(141..361)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 146..366 id H19708

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 143..264
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 143..264

id H20045

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 1..74
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100

region 4..77 id H20045

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 143..382
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 8..247

id C15772

- (ix) FEATURE:
 - (A) NAME/KEY: other

(C)	LOCATION: 157341 IDENTIFICATION METHO OTHER INFORMATION:	DD: blastn identity 98 region 1185 id H67240 est							
(B) (C)	URE: NAME/KEY: other LOCATION: 340382 IDENTIFICATION METHO OTHER INFORMATION:	DD: blastn identity 100 region 185227 id H67240 est							
(B) (C)	URE: NAME/KEY: other LOCATION: 172382 IDENTIFICATION METHO OTHER INFORMATION:	DD: blastn identity 97 region 1211 id HUM408E11B est							
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 288 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 7</pre>									
		TG GGT CCT GAT GCT GTT TCT GTG eu Gly Pro Asp Ala Val Ser Val -20 -15	49						
	Leu Gly Leu Ser Pro	GGC ACC AGC TCT GGC CGC AAC Gly Thr Ser Ser Gly Arg Asn	97						
		CTG CCG GRT YCC AGC DTC CAA Leu Pro Xaa Xaa Ser Xaa Gln 15	145						
		CCA GCC CTA CTC CTG TGG AAT Pro Ala Leu Leu Leu Trp Asn 30 35	193						
		CTG GTC CCG ACC TTG TGC CCA Leu Val Pro Thr Leu Cys Pro 45	241						
	ı Ile Leu Cys Pro Ala	CAG ACC CTG ATC CTG TGC CCA Gln Thr Leu Ile Leu Cys Pro 65	289						
	A ACC CTG TGT CCT GCC		337						

70

75

80

GTG GCA CTG TCA GCC CAG CCC TCC CTA CCG GCG AGA GTC CAG AGT
Val Ala Leu Ser Ala Gln Pro Ser Leu Pro Ala Arg Val Gln Ser
85 90 95

- (2) INFORMATION FOR SEQ ID NO: 225:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 138 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(2..139)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 135..272

id HSB82C022

est

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 10..108
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.8 seq FTSASLLLPMSTG/MP
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 225:
- ATTATTTAT ATG ATT AAC CCC TCA GTC CCT AGC AAG TCA AAT TCC CAT CCG

 Met Ile Asn Pro Ser Val Pro Ser Lys Ser Asn Ser His Pro

 -30

 -25
- TTT TTA TCT ACA GTA ATG TTC ACC TCT GCA TCA CTG CTG CTT CCC ATG

 Phe Leu Ser Thr Val Met Phe Thr Ser Ala Ser Leu Leu Leu Pro Met

 -15

 -10

 -5

TCT ACA GGC ATG CCA ACT CAA AAC TGT TTT ACC CCA AAG
Ser Thr Gly Met Pro Thr Gln Asn Cys Phe Thr Pro Lys

1 5 10

- (2) INFORMATION FOR SEQ ID NO: 226:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 406 base pairs
 - (3) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE

WO 99/065	550 PCT/IB	98/0123
	(D) TOPOLOGY: LINEAR	
(ii)	MOLECULE TYPE: CDNA	
(vi)	ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Cancerous prostate	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 138186 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 91 region 1462 id AA111755 est	
(ix)	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 83286 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.7 seq IACLAWWIGGGSG/XN	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 226:	
AAAGACTTTG	CGAASGCTGC GCTCGCGCCC GGATCCCTCA GGCGGCTGCA GGCTTCAGCC	60
TGCGCTGGTT	GGTGAAACAG AG ATG TCA GAA AAG GAG AVC AAC TTC CCG CCA 1	12

Met Ser Glu Lys Glu Xaa Asn Phe Pro Pro

160

208

256

304

352

400

406

-65

CTG CCC AAG TTC ATC CCT GTG AAG CCC TGC TTC TAC CAG AAC TTC TCC

Leu Pro Lys Phe Ile Pro Val Lys Pro Cys Phe Tyr Gln Asn Phe Ser

GAC GAG ATC CCA GTG GAG CAC CAG GTC CTG GTG AAG AGG ATC TAC CGG

Asp Glu Ile Pro Val Glu His Gln Val Leu Val Lys Arg Ile Tyr Arg -35

CTG TGG ATG TIT TAC TGC GCC ACC CTC GGC GTC AAC CTC ATT GCC TGC

Leu Trp Met Phe Tyr Cys Ala Thr Leu Gly Val Asn Leu Ile Ala Cys

CTG GCC TGG TGG ATC GGC GGA GGC TCG GGG NNB AAC TTC GGC CTG GCC

Leu Ala Tro Tro Ile Gly Gly Gly Ser Gly Xaa Asn Phe Gly Leu Ala

TTC GTG TGG CTG CTC CTG TTC ACG CCT TGC GGC TAC GTG TGC TGG TTC

Phe Val Trp Leu Leu Phe Thr Pro Cys Gly Tyr Val Cys Trp Phe

CGG CCT GTC TAC AAG GCC TTC CGA GCC GAC AGC TCC TTT AAT TTC ATG

Arg Pro Val Tyr Lys Ala Phe Arg Ala Asp Ser Ser Phe Asn Phe Met 30

-20

-55

GCG CTG

Ala Leu 40

(2) INFORMATION FOR SEQ ID NO: 227:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 347 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (68..131) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 95 region 392..455 id W22335 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 288..347 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 95 region 9..68 id H70453 (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 159..227 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.7 seq ILRLYFFLQLAHS/GY (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 227: ACGAAATGGT ATTGACATCT TGGTTGGAAC ACCTGGTCGT ATCAAAGACC ATCTGCAGAG 60 TGGCCGATTG GATCTTTCTA AACTGCGACA TGTTGTGCTT GATGAAGTGG ATCAGATGTT 120 AGATTTAGGT TTCGCTGAAC AAGTTGAAGA TATTATTC ATG AAT CCT ACA AAA CTG 176 Met Asn Pro Thr Lys Leu -20ATT CTG AAG ACA ATC CTC AGA CTT TAC TTT TTT CTG CAA CTT GCC CAC 224 Ile Leu Lys Thr Ile Leu Arg Leu Tyr Phe Phe Leu Gln Leu Ala His -15 AGT GGG TAT ACA AAG TTG CAA AAA AAA TAC ATG AAA TCC AGA TAT GAA 272 Ser Gly Tyr Thr Lys Leu Gln Lys Lys Tyr Met Lys Ser Arg Tyr Glu 5 CAG GTT GAC STT GTT GGR AAA ATG WCT CAA AAG GCT GCA ACT ACT GTG

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Gln Val Asp Leu Val Gly Lys Met Xaa Gln Lys Ala Ala Thr Thr Val 20 25 30

GRA CAT TTG GCC ATC CAG TGT CAT TGG Xaa His Leu Ala Ile Gln Cys His Trp 347

(2) INFORMATION FOR SEQ ID NO: 228:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 406 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 12..70
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96

region 1..59 id AA013305

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 197..250
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100

region 189..242 id AA013305

est

- (ix) FEATURE:.
 - (A) NAME/KEY: other
 - (B) LOCATION: 250..297
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 243..290

id AA013305

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 136..199
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 236..299

id R48472

- (ix) FEATURE:
 - (A) NAME/KEY: other

(B) LOCATION: 37..101

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 135..199 id R48472

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 38..106

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 6.7

seq SXXCFVSVPPASA/IP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 228:

AACCCGGGAC CGAGCTGGGG TCTTGGAGGA AGAGAGG ATG GCG TCG TCG AGC CCT Met Ala Ser Ser Ser Pro -20											55					
GAC Asp	TCC Ser	CCA Pro -15	TGT Cys	TCC Ser	TGS Xaa	NAC Xaa	TGC Cys -10	TTT Phe	GTC Val	TCC Ser	GTG Val	CCC Pro -5	CCG Pro	GCC Ala	TCA Ser	103
GCC Ala	ATC Ile 1	CCG Pro	GST Xaa	GTG Val	AKC Xaa 5	TTK Xaa	GCC Ala	NNH Xaa	AAC Asn	TCG Ser 10	GAC Asp	SGA Xaa	CCC Pro	CGG Arg	GAC Asp 15	151
GAG Glu	GTG Val	CAG Gln	GAG Glu	GTG Val 20	GTG Val	TTT Phe	GTC Val	CCC Pro	GCA Ala 25	GGC Gly	ACT Thr	CAC His	ACT Thr	CCT Pro 30	GGG Gly	199
AGC Ser	CGG Arg	CTC Leu	CAG Gln 35	TGC Cys	ACC Thr	TAC Tyr	ATT Ile	GAA Glu 40	GTG Val	GAA Glu	CAG Gln	GTG Val	TCG Ser 45	AAG Lys	ACG Thr	247
CAC His	GCT Ala	GTG Val 50	ATT Ile	CTG Leu	AGC Ser	CGT Arg	CCT Pro 55	TCT Ser	TGG Trp	CTA Leu	TGG Trp	GGG Gly 60	GCT Ala	GAG Glu	ATG Met	295
GGC Gly	GMV Xaa 65	ACG Thr	AGC Ser	ATG Met	GTG Val	TCT Ser 70	GCA Ala	TTG Leu	GCA Ala	ACG Thr	AGG Arg 75	CTG Leu	TGT Cys	GGA Gly	CGA Arg	343
AGG Arg 80	AGC Ser	CAG Gln	TTG Leu	GGG Gly	AGG Arg 85	GCN Ala	GKN Xaa	GCC Ala	CTS Leu	CTG Leu 90	GGC Gly	ATG Met	GAC Asp	CTA Leu	CTC Leu 95	391
	TGC Cys															406

(2) INFORMATION FOR SEQ ID NO: 229:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 308 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 128..197

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 158..227 id AA249540

est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 241..309

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 92

region 267..335 id AA249540

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(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 164..240

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 58..134 id N46699

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 128..161

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 23..56 id N46699

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement(224..309)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 104..189 id W39777

et

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 233..309

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 13..89 id AA036848

(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 233309 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 96 region 1389 id AA133513 est	
(ix)	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 171287 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.7 seq XLIAXLEPPGAMA/VR	
(xi) \$	SEQUENCE DESCRIPTION: SEQ ID NO: 229:	
CATTATTCCT :	TTTCCATCGG AAGTGGCGCT CGTGCATTCA ACTTGTTCCC GCTCATGGAA	60
CCCCTCTTTA A	AAAAGACGCA GGGCACCTGT GAGCGCAGGA GCGAGCCTAA GGCCACCCAG 1	20
CGGCAGCGCC (CGTGTCCTGG GCACTCAGCG TGCTGGGCAG AGCAGGTGCG ATG GSC Met Xaa	76
CCA GTC CTA Pro Val Leu -35	GCA GCC CTC GCC CAT GTC CTG TGC CCT TAC ATG GCT CCC Ala Ala Leu Ala His Val Leu Cys Pro Tyr Met Ala Pro -30 -25	224
GGA CTG TGC Gly Leu Cys -20	AGG GAG CCG ATA CGT TTK CTG ATA GCA VTA CTG GAA CCA Arg Glu Pro Ile Arg Xaa Leu Ile Ala Xaa Leu Glu Pro -15 -10	? 7 2
CCG GGT GCG Pro Gly Ala -5	ATG GCA GTK AGG AGA CTG CCC AGT GCC Met Ala Val Arg Arg Leu Pro Ser Ala 1 5	808
(2) INFORMAT	TION FOR SEQ ID NO: 230:	
(i) SE	EQUENCE CHARACTERISTICS: (A) LENGTH: 327 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) M	OLECULE TYPE: CDNA	
(vi) C	ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Hypertrophic prostate	
(ix) E	FEATURE: (A) NAME/KEY: other (B) LOCATION: 19327 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 99 region 1309	

id C16848

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- (A) NAME/KEY: other
- (B) LOCATION: 75..104
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 303..332

id R40385

(ix) FEATURE:

- (A) NAME/KEY: sig peptide
- (B) LOCATION: 73..207
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 6.7

seq PMLGLAAFRWIWS/RE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 230:

AAAAGCGGAC CCGCGGACGG TGGCGTTAAG GGAACGCTGA GGTCCCGCGC TCCCCGACCG

- AGGTATATCT CC ATG AAT AAC CTA AAT GAT CCC CCA AAT TGG AAT ATC CGG 111 Met Asn Asn Leu Asn Asp Pro Pro Asn Trp Asn Ile Arg -40
- CCT AAT TCC AGG GCG GAT GGT GGT GAT GGA AGC AGG TGG AAT TAT GCC 159 Pro Asn Ser Arg Ala Asp Gly Gly Asp Gly Ser Arg Trp Asn Tyr Ala -25
- CTG TTG GTT CCA ATG CTG GGA TTG GCT GCT TTT CGT TGG ATT TGG TCT 207 Leu Leu Val Pro Met Leu Gly Leu Ala Ala Phe Arg Trp Ile Trp Ser -15 -10
- AGG GAG TCC CAG AAA GAA GTA GAA AAA GAG AGA GAA GCC TAC CGT CGG 255 Arg Glu Ser Gln Lys Glu Val Glu Lys Glu Arg Glu Ala Tyr Arg Arg 1 5 10
- AGA ACT GCT GCT TTT CAA CAG GAT CTG GAA GCC AAG TAC CAC GCC ATG 303 Arg Thr Ala Ala Phe Gln Gln Asp Leu Glu Ala Lys Tyr His Ala Met 20 30

ATC TCA GAM AAT CGG CGT GCT GTC 327 Ile Ser Xaa Asn Arg Arg Ala Val

(2) INFORMATION FOR SEQ ID NO: 231:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 381 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:

WO 99/06550 214 (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate (ix) FEATURE: (A) NAME/KEY: other (3) LOCATION: complement(3..297) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 99 region 1..295 id W57719 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (37..300) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 99 region 10..273 id H04979 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (7..41) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 270..304 id H04979 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(37..295) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 99 region 7..265 id H10390 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(2..41) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 262..301 id H10390 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(142..295) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 99 region 1..154 id W42765 est (ix) FEATURE:

- (A) NAME/KEY: other
- (3) LOCATION: complement(2..141)
- (C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100 region 156..295 id W42765 est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (55..238) (C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 71..254 id R39116 est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement(255..297)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 10..52 id R39116 est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 295..351

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 6.6

seq AALCSLFFFLSLQ/EI

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 231:

ACGTTAGGGG GCCAGGGAGA TGTGACTGAG GCTGGCTTTC CACGTGAATG AGACGGGGTC 60 GGTGGAGGGT TTGGTGCTAC AGCCAGTCAG AAGATTTGCA AATGCGAACA CATTCCTGTG 120 TGAGGCACGT TACCCTTTGT CAGTTATTGT GAATATGTGT ATTTTAAGCA ATAAGATTCA 240 AGACAGAGTG GCTCTAACCA CTGTGAGAAG CCCAAATAAA AATTGATCCC AAAA ATG 297 Met CTA CTG CTC TTT CTT GCT GCA CTT TGT TCC CTC TTC TTC TTC CTC AGT 345 Leu Leu Phe Leu Ala Ala Leu Cys Ser Leu Phe Phe Leu Ser -15 -5 CTT CAG GAA ATT GCA CCT CAA GAT CCC AAA CCA GGG 381 Leu Gln Glu Ile Ala Pro Gln Asp Pro Lys Pro Gly 1

(2) INFORMATION FOR SEQ ID NO: 232:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 178 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 17..175 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 91 region 1..159 id W51023 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 42..173 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 1..132 id T61976 est (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 2..142 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.5 seq IIVCLFAFLVAHC/FL (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 232: T ATG TTA TTC CTT GGC AAG GTG CTG ATA GTC TGC AGC ACA GGT TTA GCT Met Leu Phe Leu Gly Lys Val Leu Ile Val Cys Ser Thr Gly Leu Ala -40 GGS ATT ATG CTG CTC AAC TAC CAG CAG GAC TAC ACA GTA TGG GTG CTG 97 Gly Ile Met Leu Leu Asn Tyr Gln Gln Asp Tyr Thr Val Trp Val Leu -25 CCT CTG ATC ATC GTC TGC CTC TTT GCT TTC CTA GTC GCT CAT TGC TTC 145 Pro Leu Ile Ile Val Cys Leu Phe Ala Phe Leu Val Ala His Cys Phe -10 CTG TCT ATT TAT GAA ATG GTA GTK GAT GCG AGG 178 Leu Ser Ile Tyr Glu Met Val Val Asp Ala Arg (2) INFORMATION FOR SEQ ID NO: 233:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 319 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

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(ii) MOLECULE TYPE: CDNA
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(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Cancerous prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(2..321)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 59..378 id AA045815

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 95..244
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 1..150 id R18658

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 240..321
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 147..228

id R18658

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 95..321
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 1..227

id R14615

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(2..200)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 3..201 id N95174

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement (36..197)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 8..169

id N93742

est

(ix) FEATURE:

(A) NAME/KEY: other

 (B) LOCATION: complement(244) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 166208 id N93742 est 		
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 191304 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.3</pre>		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 233:		
AAGACTCATA GAGATTAAAT GATCACTATG GTCCTTCTTC TGTTAAATGG AGG		60
GCCTATGTTG TTCTGAAGTC TTGTAATGTT TAACTTCTGA GAACTTAGAT TAG	STGGTGTG	120
ATGATAGAGT CTGTATAACG CATTGAAAAG GGTATCAGGC TTAGTTATTT ATG	CCAATAAA	180
TATTTATTGT ATG CAG GGT ATT CCT ATT TTA ACT CCT GTG ACA ACA Met Gln Gly Ile Pro Ile Leu Thr Pro Val Thr Thr -35	A CAA Gln	229
AGC ATA GCG ATT TCC ATA GTT CTA ACT GTT CAG GGT CTG CTC CT Ser Ile Ala Ile Ser Ile Val Leu Thr Val Gln Gly Leu Leu Leu -25 -15	CC CTG eu Leu -10	277
GTA CAC TCT TTT TGG TTC ACT GTA TGT ACT CCT GTT GTC TTT Val His Ser Phe Trp Phe Thr Val Cys Thr Pro Val Val Phe -5 1 5		319
(2) INFORMATION FOR SEQ ID NO: 234:		
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 360 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR		
(ii) MOLECULE TYPE: CDNA		
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate</pre>		
<pre>(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(131360) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100</pre>		

id M78402

est

219 (ix). FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (57..234) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 10..187 id H04786 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (7..43) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 201..237 id H04786 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (57..234) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 10..187 id H17078 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (7..43) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 201..237 id H17078 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (57..217) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 1..161 id HSC0UC022 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(1..43) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 175..217 id HSC0UC022 est (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 199..279 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.3

seq LFCVLLSLRPHTS/GT

WO 99/06550 PCT/IB98/01232

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 234:

ACAAGATTTT CCAACCTTGC TGGCTACTTT AGTTTGGGAC CTGTTTTTTT TCTCATTTGA TTTTGCTTGT GCAGAAAATA GTTTCCAGCA CATGGATTGA TCTGAGAGAG AATGAGGCTC 120 AGTTGTGGAT AGTCTGTTTT CTCTGAGCAT GTTGGCCAAC TAGTATCGTC AAATTATTGA 180 GTGGATCATC TCTTGGAA ATG CAG AAC TTC TGC CAC CAC TTG GCT ATT TGC Met Gln Asn Phe Cys His His Leu Ala Ile Cys ACA GTC ATC TTG TTC TGT GTC CTT TTA TCT CTC AGA CCA CAC ACA TCT Thr Val Ile Leu Phe Cys Val Leu Leu Ser Leu Arg Pro His Thr Ser -15 -10 GGA ACG CTG TGG GCA TCT TCT GCC CAT GGG CTC CAT TTG GCA CCT GCT Gly Thr Leu Trp Ala Ser Ser Ala His Gly Leu His Leu Ala Pro Ala 10 GAG CCA CAG TTG TCC TGC TGG ATG TGC TGT GCA 360 Glu Pro Gln Leu Ser Cys Trp Met Cys Cys Ala 20

(2) INFORMATION FOR SEQ ID NO: 235:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 438 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- '(ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 135..426
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 35..326 id H97426

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 92..316
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95 region 14..238

id W44834

est

- (ix) FEATURE:
 - (A) NAME/KEY: other

(B) LOCATION: 127..177 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 96 region 4..54 id R57989 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 182..211 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 93 region 62..91 id R57989 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (287..316) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 96 region 385..414 id N93806 (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 34..225 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.3 seg VLMRLVASAYSIA/OK (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 235: AAGTTTCCCG CATGCTCAGT AGCTGAGGTA GGG ATG CCA TCC TTC TCA AAA GAC Met Pro Ser Phe Ser Lys Asp -60 TTA TTG ACA GTG CCA AAG CTC GGT ACT GGA CAC VMC GRR GGR MCT GGG 102 Leu Leu Thr Val Pro Lys Leu Gly Thr Gly His Xaa Xaa Gly Xaa Gly TCC TAC GAT RAC GCG CTT KTG CTC CTC CTG AAG TGT CTT TGG TCC AAC 150 Ser Tyr Asp Xaa Ala Leu Xaa Leu Leu Leu Lys Cys Leu Trp Ser Asn GTT GTT CCA GAG TGT ACC ATG GCT TCC AGT AAC ACT GTG TTG ATG CGG Val Val Pro Glu Cys Thr Met Ala Ser Ser Asn Thr Val Leu Met Arq -20-15 TTG GTA GCC TCC GCA TAT TCT ATT GCT CAA AAG GCA GGA ATG ATA GTC 246 Leu Val Ala Ser Ala Tyr Ser Ile Ala Gln Lys Ala Gly Met Ile Val AGA CGT GTT ATT GCT GAA GGA GAC CTG GGT ATT GTG GAG AAG ACC TGT 294 Arg Arg Val Ile Ala Glu Gly Asp Leu Gly Ile Val Glu Lys Thr Cys

GCA ACA GAC CTG CAG ACC AAA GCT GAC CGA TTG GCA CAG ATG AGC ATA

Ala Thr Asp Leu Gln Thr Lys Ala Asp Arg Leu Ala Gln Met Ser Ile

342

WO 99/06550 PCT/IB98/01232

25

30

35

TGT TCT TCA TTG GYM BGG AAA TTC CCC AAA CTC RNR ATT ATA GGG GAA

Cys Ser Ser Leu Xaa Xaa Lys Phe Pro Lys Leu Xaa Ile Ile Gly Glu

40 50 55

GAG GAT CTG CCT TCT GAG GAA GTG GAT CAA GAG CTG ATT GAA GAC AGK
Glu Asp Leu Pro Ser Glu Glu Val Asp Gln Glu Leu Ile Glu Asp Xaa
60 65 70

(2) INFORMATION FOR SEQ ID NO: 236:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 310 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 7..113
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 15..121

id W04921

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 114..220
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 121..227

id W04921

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 221..310
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 93

region 227..316

id W04921

est

- (ix) FEATURE:
 - .(A) NAME/KEY: other
 - (B) LOCATION: complement (114..213)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 260..359

id N70602

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(32..113)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 359..440

id N70602

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement (261..311)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 92

region 164..214

id N70602

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement (213..259)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 95

region 215..261

id N70602

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 114..194
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 59..139

id W70167

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 238..311
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 183..256

id W70167

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 55..113
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 1..59

id W70167

est.

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 193..236
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 95

region 139..182

id W70167 est

(ix) FEATURE:

- (A) NAME/KEY: other
 (B) LOCATION: 221..311
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 91

region 165..255

id W37690

est

(ix) FEATURE:

- (A) NAME/KEY: other(B) LOCATION: 114..187
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 58..131

id W37690

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 56..113
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 1..58

id W37690

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 185..220
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 130..165

id W37690

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 227..289
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 6.2

seq LEMLXAFASHIXA/RD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 236:

ATGGCAGCTT CCTTGGCTCG GCTTGGTCTG CGGCCTGTCA AACAGGTTCG GGTTCAGTTC 60

TGTCCCTTCG AGAAAAACGT GGAATCGACG AGGACCTTCV TSCAGACGGT GAGGCMGTGA 120

GAAGGTCCGC TCCACTAATC TCAACTGCTC AGTGATTGCG GACGTGASGC ATGACGGCTC 180

CGAGCCCTGC GTGGACGTGC TGTTCGGAGA CGGGCATCGC CTGATT ATG CGC GGC 235

Met Arg Gly

-20

GCT CAT CTC ACC GCT CTG GAA ATG CTC ANM GCC TTC GCC TCC CAC ATM
Ala His Leu Thr Ala Leu Glu Met Leu Xaa Ala Phe Ala Ser His Ile

PCT/IB98/01232 WO 99/06550 225

> -15 -10 -5

HGG GCC AGG GAC GCG GCG GGC AGC GGG Xaa Ala Arg Asp Ala Ala Gly Ser Gly 1

310

(2) INFORMATION FOR SEQ ID NO: 237:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 429 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 321..431
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 186..296 id AA043558

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 218..299
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 93

region 83..164 id AA043558

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 173..230
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100

region 39..96

id AA043558

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 131..299
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 57..225

id N50523

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 321..431

(C) IDENTIFICATION METHOD: blastn
(D) OTHER INFORMATION: identity 98
region 247..357
id N50523
est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement (45..115)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 1..71 id N50523 est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement (321..431)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 289..399 id AA115605

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(217..318)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 403..504 id AA115605

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(166..231)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 92

region 491..556 id AA115605

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 172..318
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 36..182 id AA115129

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 321..431
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 186..296

id AA115129

est

(ix) FEATURE:

(B) L(C) I	AME/KEY: other OCATION: 174318 DENTIFICATION MET THER INFORMATION:	***	
(ix) FEATUR	E: AME/KEY: other		
• •	OCATION: 325431		
	DENTIFICATION MET	HOD: blastn	
• •	THER INFORMATION:		
(ix) FEATUR	E:		
	AME/KEY: sig_pept	ide	
	OCATION: 7423	HOD: Von Heijne matrix	
	THER INFORMATION:		
,-, -		seq FGLLHQLSQCVTS/LE	
(xi) SEQUEN	CE DESCRIPTION: S	EQ ID NO: 237:	
		ATT ACC CTC TTT CTC ACC AGC	48
Met Glu Va	_	Ile Thr Leu Phe Leu Thr Ser	
	-135	-130	
GCC AGC AGC CCT G	TG GTG GCG ACG AC	G ATG GAC CAG GAG CCA GTG GGC	96
		r Met Asp Gln Glu Pro Val Gly	
-125	-120	-115 -110	
GGT GTG GAA CGA G	GA GAA GCC GTC GC	A GCC TCG GGA RCT GCG GCC GCC	144
		a Ala Ser Gly Xaa Ala Ala Ala	144
-	105	-100 -95	
ברר רבת ששר הכה ה	מא דרד כרא כככ לא	G ATG AGT AAC GAA AGA GGC TTT	100
		n Met Ser Asn Glu Arg Gly Phe	192
90	-8		
		A AAA AAG AAG AAA GTC CCA AGG y Lys Lys Lys Val Pro Arg	240
-75	-70	y Lys Lys Lys Lys Vai Pro Arg -	
. •	. •	03	
AGA GTC ATC CAC T	TT GTT AGT GGT GA	A ACA ATG GAA GAA TAT AGC ACA	288
	he Val Ser Gly Gl? -55	u Thr Met Glu Glu Tyr Ser Thr	
-60	-55	-50	
GAT GAA GAC GAH G	TT GAT GGC CTG GA	G AAG NNG ATG TTT TGC CTA CTG	336
		u Lys Xaa Met Phe Cys Leu Leu	
-45	-40	-35 -30	
TTG ATC CGR CAA A	אר דדא ככד ההה הז	'C CCT ACT TAT GGT TTT ACA TGC	301
		of Pro Thr Tyr Gly Phe Thr Cys	384
	-25	-20 -15	
	O O		
FIC GGG CTG CTA C	AI CAA CTC TCT CA	NG TGT GTG ACT TCC TTG GAG	429

Phe Gly Leu Leu His Gln Leu Ser Gln Cys Val Thr Ser Leu Glu

-5

228

-10

1

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(2) INFORMATION FOR SEQ ID NO: 238:
```

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 321 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Cancerous prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 102..322
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 31..251

id T34679

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 176..322
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 104..250

id N34677

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 93..170
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 91

region 21..98

id N34677

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 180..312
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 203..335

id N32531

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 180..312
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 202..334

id N36824

	est
(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 102170 (C) IDENTIFICATION METHORMATION:	HOD: blastn identity 97 region 2896 id N36824 est
<pre>(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 175312 (C) IDENTIFICATION METH (D) OTHER INFORMATION:</pre>	
(ix) FEATURE: (A) NAME/KEY: sig_pepti (B) LOCATION: 151279 (C) IDENTIFICATION METH (D) OTHER INFORMATION:	HOD: Von Heijne matrix
(xi) SEQUENCE DESCRIPTION: SE	EQ ID NO: 238:
AACTCTCGTG CCAAGCATGT CTCTCCAAAT GO	GCTGCTCTC TGGCGTTCCT CACACTCCCC 60
CTGAAGTTCA TCTAAGATCT TCATTCTTCA WA	AGGCGGAAG CCCGGCTCGC TGCAAAACGG 120
GCGGCCCGCG CGGAGGCTCG CGAGATCCGC A	FG AAG GAG CTG GAG CGG CAG 174 et Lys Glu Leu Glu Arg Gln Gln -40
AAG GAG GTA GAA GAG AGA CCA GAA AA Lys Glu Val Glu Glu Arg Pro Glu Ly -35	
CGT AAC ATG CCG GGC CTG TCT GCA GCC Arg Asn Met Pro Gly Leu Ser Ala Ala -15	
ACT TCC TCT CGG AGA GGC AGC GGA GA Thr Ser Ser Arg Arg Gly Ser Gly As 1 5	
GAG Glu	321

(2) INFORMATION FOR SEQ ID NO: 239:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 401 base pairs
(B) TYPE: NUCLEIC ACID
(C) STRANDEDNESS: DOUBLE

WO 99/06550 PCT/IB98/01232

230 (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 270..403 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 92 region 199..332 id AA125491 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 70..135 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 93 region 1..66 id AA125491 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (27..135) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 89..197 id HSB72F052 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(135..223) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 2..90 id HSB72F052 est (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 126..188 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6.1 seq VLVILCIVTVCVT/IV (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 239: ACCGGAGAAA AAATGGTTCA TGGAGCCTGC GGTTATTGTT TGCCTGGGTG GAATTTTACC 60 TTTTGGTTCA ATCTTTATTG AAATGTATTT CATCTTCACG TCTTTCTGGG CATATAAGAT CTATT ATG TCT ATG GGC TTC ATG ATG CTG GTG CTG GTT ATC CTG TGC ATT 170

Met Ser Met Gly Phe Met Met Leu Val Leu Val Ile Leu Cys Ile

-15

-20

GTG ACT GTC TGT GTG ACT ATT GTG TGC ACA TAT TTT CTA CTA AAT GCA
Val Thr Val Cys Val Thr lle Val Cys Thr Tyr Phe Leu Leu Asn Ala
10

GAA GAT TAC AGG TGG CAA TGG ACA AGT TTT CTC TCT GCT GCA TCA ACT
Glu Asp Tyr Arg Trp Gln Trp Thr Ser Phe Leu Ser Ala Ala Ser Thr
15

GCA ATC TAT GTT TAC ATG TAT TCC TTT TAC TAC TAT TTT TTC AAA ACA
Ala lle Tyr Val Tyr Met Tyr Ser Phe Tyr Tyr Tyr Phe Phe Lys Thr
30

AAG ATG TAT GGC TTA TTT CAA ACA TCA TTT TAC TTT GGA TAT ATG GCG
Lys Met Tyr Gly Leu Phe Gln Thr Ser Phe Tyr Phe Gly Tyr Met Ala
45

GTA TTT AGC ACA GCC TTG GGG ATA ATG TGT GGA GCG ATT
Val Phe Ser Thr Ala Leu Gly Ile Met Cys Gly Ala Ile
60

70

(2) INFORMATION FOR SEQ ID NO: 240:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 466 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 153..397
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 131..375

id W56159 est

- (ix) FEATURE:
 (A) NAME/KEY: other
 - (B) LOCATION: 19..139
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100

region 1..121

id W56159

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 153..467
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95 region 303..617

id HSZ78368 est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 60..139
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 214..293

id HSZ78368

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 153..374
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 80..301

id AA026570

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 70..139
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 1..70

id AA026570

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 372..405
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 300..333

id AA026570

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 155..467
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 90

region 83..395

id AA109961

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 88..139
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 92

region 20..71

id AA109961

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 153..363
- (C) IDENTIFICATION METHOD: blastn

WO 99/06550 PCT/IB98/01232

(D) OTHER INFORMATION: identity 96 region 274..484 id AA046907 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 60..139 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 185..264 id AA046907 est (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 128..337 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 6 seq LLFPLTLVRSFWS/DM (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 240: AACGCTTGCG ATGGTTGAAT TCCCCTCCTC ACGCCAGCCT AGGAGAAGAA GTTCGTAGTC CCAGAGGAAG AGGAGTTGTA CGCATGTCAG AGAGGTTGCA GGCTGTTTTC AATTTGTCAG 120 TTTGTGG ATG ATG GAA TTG GRM CTW AAA AKC GRA ACT AAA KKG GAA TGK 169 Met Met Glu Leu Xaa Leu Lys Xaa Xaa Thr Lys Xaa Glu Xaa -65 GAA TCT GCA TGT ACA GAA GCA TAT TCC CAA TCT GAT GAG CAA TAT GCT 217 Glu Ser Ala Cys Thr Glu Ala Tyr Ser Gln Ser Asp Glu Gln Tyr Ala -55 -50 TGC CAT CTT GGT TGC CAG AAT CAG CTG CCA TTC GCT GAA CTG AGA CAA 265 Cys His Leu Gly Cys Gln Asn Gln Leu Pro Phe Ala Glu Leu Arg Gln -40 -35 -30 GAA CAA CTT ATG TCC CTG ATG CCA AAA ATG CAC CTA CTC TTT CCT CTA 313 Glu Gln Leu Met Ser Leu Met Pro Lys Met His Leu Leu Phe Pro Leu -20 -15 ACT CTG GTG AGG TCA TTC TGG AGT GAC ATG ATG GAC TCC GCA CAG AGC 361 Thr Leu Val Arg Ser Phe Trp Ser Asp Met Met Asp Ser Ala Gln Ser TTC AKA ACC TCT TCA TGG ACT TTT TAT CTT CAA GCC GAT GMC GGM MAA Phe Xaa Thr Ser Ser Trp Thr Phe Tyr Leu Gln Ala Asp Xaa Gly Xaa 10 15 ATA GTT ATA TKC CAG TCT AAG CCA GAA ATC CAG TAC GCA CCA CAT TTG 457 Ile Val Ile Xaa Gln Ser Lys Pro Glu Ile Gln Tyr Ala Pro His Leu 25 35 GAG CAG GAG 466 Glu Gln Glu

(2) INFORMATION FOR SEQ ID NO: 241:

```
(i) SEQUENCE CHARACTERISTICS:
```

- (A) LENGTH: 81 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 18..81
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 62..125 id AA092155 est
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(18..81)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 68..131 id AA128307 est
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(18..81)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 68..131 id N99068 est

- '(ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(18..81)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 68..131 id AA039944 est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(18..81)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 68..131 id AA128099
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide

(B) LOCATION: 1..72

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 6

seq GLILLFASHLINQ/FS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 241:

ATG GTT TCC AAT GCT TCR GAG ACT TCC TGC CTA GGC CTC ATC CTC CTC

Met Val Ser Asn Ala Ser Glu Thr Ser Cys Leu Gly Leu Ile Leu Leu

-20

-15

-10

TTT GCC AGT CAC CTG ATT AAC CAA TTC TCC AGC
Phe Ala Ser His Leu Ile Asn Gln Phe Ser Ser
-5

(2) INFORMATION FOR SEQ ID NO: 242:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 373 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Hypertrophic prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 29..302

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99 region 1..274

id H18735

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 143..302

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 116..275

id T80360

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 79..143

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 92

region 51..115

id T80360

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 29..69

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100 region 1..41

id T80360

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 66..302
- (C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 1..237 id AA137006

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 301..336
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 412..447 id AA137006

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 65..302
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 2..239 id HSC2CA081

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 64..224
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 1..161

id T36290

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 223..302
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 161..240

id T36290

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 2..220
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 6

seq LIVFISVCTALLA/EG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 242:

A A	rg co	CC CC ro Ai	rg L	AG CO ys Ai 70	GG AM	AG TO	GC GA	sp L	IT Co eu A: 65	GG GG	CT G' la Va	TC AG	rg Va	TT GO al Gi 60	GT CTG ly Leu	49
TTA Leu	CTC Leu	GGT Gly -55	GGT Gly	GGC Gly	GGA Gly	GTC Val	TAC Tyr -50	GGA Gly	AGC Ser	CGT Arg	TTT Phe	CGC Arg -45	TTC Phe	ACT Thr	TTT Phe	97
CCT Pro	GGC Gly -40	TGT Cys	AGA Arg	GCG Ala	CTT Leu	TCC Ser -35	CCC Pro	TGG Trp	CGG Arg	GTG Val	AGA Arg -30	VTG Xaa	CAG Gln	AGA Arg	CGA Arg	145
AGG Arg -25	TGC Cys	GAG Glu	ATG Met	AGC Ser	ACT Thr -20	ATG Met	TTC Phe	GCG Ala	GAC Asp	ACT Thr -15	CTC Leu	CTC Leu	ATC Ile	GTT Val	TTT Phe -10	193
ATC Ile	TCT Ser	GTG Val	TGC Cys	ACG Thr ~5	GCT Ala	CTG Leu	CTC Leu	GCA Ala	GAG Glu 1	GGC Gly	ATA Ile	ACC Thr	TGG Trp 5	GTC Val	CTG Leu	241
GTT Val	TAC Tyr	AGG Arg 10	ACA Thr	GAC Asp	AAG Lys	TAC Tyr	AAG Lys 15	AGA Arg	CTG Leu	AAG Lys	GCA Ala	GAA Glu 20	GTG Val	GAA Glu	AAA Lys	289
												ASC Xaa				337
	CCC Pro															373

(2) INFORMATION FOR SEQ ID NO: 243:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 447 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 159..307
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95 region 121..269

id W31320 est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 37..121

(C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 95

region 1..85 id W31320 est

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 320..380

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 282..342

id W31320

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 114..165

(C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 94

region 77..128

id W31320

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 400..443

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 364..407

id W31320

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 154..307

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 2..155

id T27259

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 320..443

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 168..291

id T27259

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 192..307

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93 region 108..223

100..22

id AA157646

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 64..95

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 1..32 id AA157646

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 320..443

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 123..246 id AA182962

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 198..307

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 1..110 id AA182962

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 243..307

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 189..253

id T71690

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 181..235

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 125..179

id T71690

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 114..164

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 94

region 58..108

id T71690

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 130..198

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.9

seq LGAAALALLLANT/DV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 243:

CCC	CGCC	CCT	GGGA	CCCT	CC G	GGCC	GGGC	G GT	TTG6	cccc	TTA	GCGC	CCG	GGCG	TCGGG	G 60
															CAAGT	
TTC	TTCA	CC A	TG G et G	GG A ly M	et T	GG T rp S 20	CC A	TT G le G	GT G	la G	GA G ly A 15	CC C la L	TG G eu G	GG G ly A	CT GC la Al -1	a
GCC Ala	TTG Leu	GCA Ala	TTG Leu	CTG Leu -5	CTT Leu	GCC Ala	AAC Asn	ACA Thr	GAC Asp	GTG Val	TTT Phe	CTG Leu	TCC Ser 5	AAG Lys	CCC Pro	219
CAG Gln	AAA Lys	GCG Ala 10	GCC Ala	CTG Leu	GAG Glu	TAC Tyr	CTG Leu 15	GAG Glu	GAT Asp	ATA Ile	GAC Asp	CTG Leu 20	AAA Lys	ACA Thr	CTG Leu	267
GAG Glu	AAG Lys 25	GAA Glu	CCA Pro	AGG Arg	ACT Thr	TTC Phe 30	AAA Lys	GCA Ala	AAG Lys	GAG Glu	CTA Leu 35	TGG Trp	GAA Glu	AAA Lys	AAT Asn	315
GGA Gly 40	GCT Ala	GTG Val	ATT Ile	ATG Met	GCC Ala 45	GTG Val	CGG Arg	AGG Arg	CCA Pro	GGC Gly 50	TGT Cys	TTC Phe	CTC Leu	TGT Cys	CGA Arg 55	363
GAG Glu	GAA Glu	GCT Ala	GCG Ala	GAT Asp 60	CTG Leu	TCC Ser	TCC Ser	CTG Leu	AAA Lys 65	AGC Ser	ATG Met	TTG Leu	GAC Asp	CAG Gln 70	CTG Leu	411
GGC Gly	GTC Val	CCC Pro	CTC Leu 75	TAT Tyr	GCA Ala	GTG Val	GTA Val	AAG Lys 80	GAG Glu	CAA Gln	CGG Arg					447

(2) INFORMATION FOR SEQ ID NO: 244:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 428 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 1..382
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 13..394 id C17481 est
- (ix) FEATURE:
 - (A) NAME/KEY: other

(B) LOCATION: 379..424

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 390..435

id C17481 est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 68..258

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 72..262 id T46941

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 1..67

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 6..72 id T46941

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (149..271)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 92

region 1..123 id R75331

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 257..430

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 42..215

id W95977

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 278..430

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 5..157 id R57521

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 255..347

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.9

seq LPLLLVANAGTAA/VG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 244:

ATGAAAATGG	GTGTGCTTAT	TTCCACGAAG	AGGAAAGAGA	AGGACTTGCA	AAGATATGTA	60
GGCTTGCCAT	TCATTCTCGA	TATGAAGACT	TCGTAGTGGA	TGGCTTCAAT	GTGTTATATA	120
				TGGCCTGGGC		180
				ACCCTGTAAC		240
GATCCCAGCA	TCAG ATG GA Met As	p Val Ala :	TTC CTG GAG Phe Leu Glu -25	ADA CTG ATT Xaa Leu Ile	AAA GAT Lys Asp -20	290
GAT ATA GAG Asp Ile Glu	G CGA GGA AG Arg Gly Ar -15	A CTG CCC (g Leu Pro 1	CTG TTG CTT Leu Leu Leu -10	GTC GCA AAT Val Ala Asn	GCA GGA Ala Gly -5	338
ACG GCA GCA Thr Ala Ala	GTA GGA CA Val Gly Hi	C ACA GAC A 5 Thr Asp I 5	AAG ATT GGG Lys Ile Gly	AGA TTG AAA Arg Leu Lys 10	GAA CTC Glu Leu	386
TGT GAG CAG Cys Glu Gln 15	TAT GGC ATA	A TGG CTT (Trp Leu E 20	CAT GTG GAG His Val Glu	GGT GTG AAT Gly Val Asn 25		428

(2) INFORMATION FOR SEQ ID NO: 245:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 233 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 1..230
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99 region 3..232 id HSC1WH101

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 102..230
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 41..169 id R12437 est
- (ix) FEATURE:
 - (A) NAME/KEY: other

(B) LOCATION: 63104		
(C) IDENTIFICATION METH		
(D) OTHER INFORMATION:	identity 100	
	region 142 id R12437	
	est	
(ix) FEATURE:		
(A) NAME/KEY: other		
(B) LOCATION: 63230		
(C) IDENTIFICATION METHO		
(D) OTHER INFORMATION:	region 1168	
	id R13448	
	est	
(ix) FEATURE:		
(A) NAME/KEY: other		
(B) LOCATION: 165212 (C) IDENTIFICATION METHO	OD: blastn	
	identity 100	
(.,	region 3683	
	id T69236	
	est	
(in) PRAMURE.		
<pre>(ix) FEATURE:</pre>	da	
(B) LOCATION: 180227	ue	
(C) IDENTIFICATION METHO	OD: Von Heijne matrix	
(D) OTHER INFORMATION:		
	seq LFNLLWLALACSP/VW	
() CEQUENCE DESCRIPTION. CE	O TD NO. 245	
(xi) SEQUENCE DESCRIPTION: SE	Q 1D NO: 245:	
GTTTGTGGCC GTCCGGCCTC CCTGACATGC AG	ATTTCCAC CCAGAAGACA GAGAAGGAGC	60
CAGTGGTCAT GGAATGGGCT GGGGTCAAAG AC	TGGGTGCC TGGGAGCTGA GGCAGCCACC	120
GTTTCAGCCT GGCCAGCCCT CTGGACCCCG AG		170
- Control Control Control Ad	GIIGGACC CIACIGIGAC ACACCIACC	179
ATG CGG ACA CTC TTC AAC CTC CTC TGG	CTT GCC CTG GCC TGC AGC CCT	227
Met Arg Thr Leu Phe Asn Leu Leu Trp	Leu Ala Leu Ala Cys Ser Pro	
-15 -10	- 5	
GTT TGG	•	
Val Trp		233
1		
(2) THEODWARTON FOR ORD TO NO. OLG		
(2) INFORMATION FOR SEQ ID NO: 246:		
(i) SEQUENCE CHARACTERISTICS:		
(A) LENGTH: 330 base pa	irs	
(B) TYPE: NUCLEIC ACID		
(C) STRANDEDNESS: DOUBL	E	
()) TOPOLOGY: LINEAR		
(ii) MOLECULE TYPE: CDNA		

(vi)	ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 178331 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 118271 id R60406 est
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 178316 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 94 region 57195 id N78477 est
(ix) I	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 214312 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.8 seq FICLQWALPHSEA/GD
(xi) S	SEQUENCE DESCRIPTION: SEQ ID NO: 246:
AAAGGCAGGA C	CTGACGCAGA ATGACAACGG CAACACGACA AGAAGTCCTT GGCCTCTACC 60
GCAGCATTTT C	CAGGCTTGCG AGGAAATGGC AGGCGACATC AGGGCAGATG GAAGACACCA 120
	ACAGTACATA CTAAATGAAG CCAGAACGCT GTTCCGGAAA AACAAAAATC 180
TCACGGACAC A	AGACCTAATT AAACAGTGTA TAG ATG AAT GCA CAG CCA GGA TTG 234 Met Asn Ala Gln Pro Gly Leu -30
AWA TTG GAC Xaa Leu Asp -25	TGC ATT ACA AGA TTC CTT ACC CAN GGC CAA TTC ATC TGC Cys Ile Thr Arg Phe Leu Thr Xaa Gly Gln Phe Ile Cys -20 -15
CTC CAA TGG Leu Gln Trp -10	GCC TTA CCC CAC TCC GAG GCC GGG GAC TTC GAA GCC AAG Ala Leu Pro His Ser Glu Ala Gly Asp Phe Glu Ala Lys -5 1 5
(2) INFORMAT	ION FOR SEQ ID NO: 247:
	QUENCE CHARACTERISTICS: (A) LENGTH: 353 base pairs

- (B) TYPE: NUCLEIC ACID
 (C) STRANDEDNESS: DOUBLE
 (D) TOPOLOGY: LINEAR

- (ii) MOLECULE TYPE: CDNA

WO 99/06550 245

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement(230..352)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 32..154

id W60134

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (78..189)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 195..306

id W60134

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (9..87)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 91

region 298..376

id W60134

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (176..352)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 57..233

id H64097

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (57..189)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 219..351

id H64097

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (84..352)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 57..325

id W00624

(ix) FEATURE:

(A) NAME/KEY: other

	(B) LOCATION: compleme (C) IDENTIFICATION MET (D) OTHER INFORMATION:	HOD: blastn	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (C) IDENTIFICATION MET (D) OTHER INFORMATION:	nt(1168) HOD: blastn identity 100 region 156323 id W67127 est	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (C) IDENTIFICATION METR (D) OTHER INFORMATION:	HOD: blastn	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (C) IDENTIFICATION METH (D) OTHER INFORMATION:	IOD: blastn	
(ix) -	FEATURE: (A) NAME/KEY: other (B) LOCATION: complemen (C) IDENTIFICATION METH (D) OTHER INFORMATION:	t(2364) OD: blastn identity 92 region 347388 id H10776 est	
(ix)	FEATURE: (A) NAME/KEY: sig_pepti (B) LOCATION: 120326 (C) IDENTIFICATION METH (D) OTHER INFORMATION:		
(xi)	SEQUENCE DESCRIPTION: SE	Q ID NO: 247:	
		CATTTTTA AAATGAAGTG TTGTTGCCTT	60
	•	GCTAAACT TTGCCTCCAG TGGTCAAAG	119
ATG GGA AAA Met Gly Lys	GAG TGG GGT TGG CAG GAG Glu Trp Gly Trp Gln Glu -65	ATG GAA AAC GGA GGT GCC GCC Met Glu Asn Gly Gly Ala Ala -60 -55	167

CCA GCA TGG GGG GCA GGT CCC CCA GTC CAC CCT GCC CCT CCC CCT GTG 215 Pro Ala Trp Gly Ala Gly Pro Pro Val His Pro Ala Pro Pro Pro Val -45 GAG AAG ACG CTT AGT TGG GGG TGT GGG TTT GGG CTC CAT TCT GGA TTC 263 Glu Lys Thr Leu Ser Trp Gly Cys Gly Phe Gly Leu His Ser Gly Phe -35 GGC GGT TCC GGG GGA GGG GTG GGT CTG TGC CGA TTA CTC TGT CTT GTA 311 Gly Gly Ser Gly Gly Val Gly Leu Cys Arg Leu Cys Leu Val -20 -15 CGT TTG TTC TGC TGC TCT TCA ATA TTG TAT CAA CGC CAG AAG 353 Arg Leu Phe Cys Cys Ser Ser Ile Leu Tyr Gln Arg Gln Lys

(2) INFORMATION FOR SEQ ID NO: 248:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 108 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (3) LOCATION: 22..71
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 94 region 1..50 id R82719 est
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 19..62
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 93

region 1..44 id AA069083

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 20..52
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 2..34

id R29193

- (ix) FEATURE:
 - (A) NAME/KEY: other

(B) LOCATION: 23..52

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 10..39 id AA158081

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 10..96

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.7

seq AALLLTATVRLSA/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 248:

AAGTCCAAC ATG GCG GCG CCC AGC GGA GGG TGG AAC GGC GTC GGC GCG AGC

Met Ala Ala Pro Ser Gly Gly Trp Asn Gly Val Gly Ala Ser

-25

TTG TGG GCC GCG CTG CTC CTC ACT GCC ACA GTC AGA CTT TCA GCT TCT
Leu Trp Ala Ala Leu Leu Thr Ala Thr Val Arg Leu Ser Ala Ser
-15 -5 1

CCC GGC CCA Pro Gly Pro

108

(2) INFORMATION FOR SEQ ID NO: 249:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 393 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Cancerous prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 7..165

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100 region 1..159 id R24141

est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 178..264

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97 region 173..259

id R24141

	(3	ix) i	(B) (C)	NAME LOCA IDEN	ATION NTIF	(: ot N: 25 ICATI	82 ON N	1ETHC	ider regi	itity	y 95 254	. 295			·
	į)	ix) i	(B) (C)	NAME LOCA I DEN	ATION NTIFI	(: ot N: 23 ICATI NFORM	03 ON M	1ETHC	ider regi	tity	/ 95 L12	20			
	(i	lx) I	(B) (C)	NAME LOCA I DEN	ATION NTIFI	(: si N: 4. ICATI NFORM	.147 ON M	r METHO	D: V	e 5.	7		itrix		
	()	(i) \$	SEQUE	ENCE	DESC	CRIPT	: NOI	: SE(Q ID	NO:	249:	:			
ATC						GGG Gly									48
						CGA Arg									96
						CTG Leu									144
						TTC Phe									192
						AGC Ser									240
						GGG Gly									288
						GTG Val									336

GAC CTG GAG CGG ACA ACG GCT CCC TGG ACG GCC CTA CTA CAT GTC CAA Asp Lau Glu Arg Thr Thr Ala Pro Trp Thr Ala Leu Leu His Val Gln 65

384

GAG CTT CTA Glu Leu Leu 80

393

(c) intollimitation for Sec 10 NO: 750	(2)	INFORMATION	FOR	SEO	ID	NO:	250
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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 363 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 222..265
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 220..263

id N89186 est

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 76..348
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.7

seq SVLELIVASVCQS/HI

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 250:

GCTACTTTCT TTTTCAGTCT TTCGGTGCGG AGAAGGGGAG GAGGCGGGCA GAGGTCTGAA 60

AAAATCGAAT GCCTT ATG GAA AGG AAC TGC AAG GGT TCC TTT GGG GTG ATC

Met Glu Arg Asn Cys Lys Gly Ser Phe Gly Val Ile

-90 -85 -80

AAA GAG GGA GAC ACA GAC ACA GRR GAG ACA AAG GCA AGG AGG ACT GTC
Lys Glu Gly Asp Thr Asp Thr Xaa Glu Thr Lys Ala Arg Arg Thr Val

-75

-70

-70

TGG GAG CCA CGC GGG CGA TAC AGT TTC CGA GRM ACG CCG CGT CCC GCC
Trp Glu Pro Arg Gly Arg Tyr Ser Phe Arg Xaa Thr Pro Arg Pro Ala
-60
-55

TAT CCT GTT GAA CAG TGC GGA TTT GCG AGG CGC GCC CTG GAG CTG CTA

Tyr Pro Val Glu Gln Cys Gly Phe Ala Arg Arg Ala Leu Glu Leu Leu

-45

-40

-35

GAG ATC CGG AAG CAC AGC CCC GAG GTG TGC GAA CCA CCA AAC ATC CCA
Glu Ile Arg Lys His Ser Pro Glu Val Cys Glu Pro Pro Asn Ile Pro
-30 -25 -20

WO 99/06550 251 GTT ACC AGT GTC CTT GAA TTG ATA GTG GCT TCT GTT TGT CAG TCT CAT 351 Val Thr Ser Val Leu Glu Leu Ile Val Ala Ser Val Cys Gln Ser His -15 ATA AGA ACT ACT 363 Ile Arg Thr Thr (2) INFORMATION FOR SEO ID NO: 251: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 293 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 22..264 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 1..243 id AA211459 est (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 15..212 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.7 seq LYMLAEALPVSHG/AH (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 251: GTGAAGATGA AGCC ATG TTT GTA GAA TAT AGA AAA CAA CTG AAG TTA CTG 50 Met Phe Val Glu Tyr Arg Lys Gln Leu Lys Leu Leu TTG GAC AGG CTT GCT CAA GTT TCA CCA GAG TTA CTA CTG GCC TCT GTT Leu Asp Arg Leu Ala Gln Val Ser Pro Glu Leu Leu Leu Ala Ser Val -45 CGC AGA GTT TTT AGT TCT ACA CTG CAG AAT TGG CAG ACT ACA CGG TTT 146 Arg Arg Val Phe Ser Ser Thr Leu Gln Asn Trp Gln Thr Thr Arg Phe

-30

194

ATG GAA GTT GAA GTA GCA ATA AGA TTG CTG TAT ATG TTG GCA GAA GCT

Met Glu Val Glu Val Ala Ile Arg Leu Leu Tyr Met Leu Ala Glu Ala -15

CTT CCA GTA TCT CAT GGT GCT CAC TTC TCA GGT GAT GTT TCA AAA GCT Leu Pro Val Ser His Gly Ala His Phe Ser Gly Asp Val Ser Lys Ala

	WO 99/06550 252													PCT/IB98/01232		
	-5					1				5					10	
AGT Ser	GCT Ala	TTG Leu	CAG Gln	GAT Asp 15	ATG Met	ATG Met	CGA Arg	ACT Thr	CTG Leu 20	GTA Val	ACA Thr	TCA Ser	GGA Gly	GTC Val 25	AGC Ser	290
GGG Gly																293
(2)	INFO	RMAT	CION	FOR	SEQ	ID 1	10: 2	252:								
	(i	.) SE	QUEN	ICE C	HARA	CTEF	RISTI	cs:								

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(A) LENGTH: 394 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: DOUBLE(D) TOPOLOGY: LINEAR

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 155..187

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93

region 95..127 id H83489

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 326..388

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.7

seq IIFLIQWHGSVFQ/EF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 252:

AAGTCCCTGT ACAGGGTTTC TGACCTGTGG TAAAAACAGA ATGTCACTTT CTGACAGGCA	60
CAGTACCCC AGGATAAACT TGGAACCTCG AGAGGAAATT CACGAAACTC GTGGGGGCAG	120
GGGTCACAAG GTGCTTGGTG GGGGARAASC TGGAAGACAT ATTGTCCAGG AGAAGGAATG	180
TCACAAGGAA CTGACAAAAT CAAGTCACGG CGCCTACAAA GATGAGGGGC AGATTCTGGC	240
TGCCTTTTAA TTTCGTCCTT CACCTGATAT CTGTGCCAGA GAATGATAAA AATCATAATA	300
AAGGRAATAG YGGAAGAGGA GACTT ATG TTA CTG GGG ACA TCT AAC ATA ATT Met Leu Leu Gly Thr Ser Asn Ile Ile -20 -15	352
ATT TTC CTG ATT CAG TGG CAT GGT TCA GTC TTC CAG GAG TTC Ile Phe Leu Ile Gln Trp His Gly Ser Val Phe Gln Glu Phe -10	394

(2) INFORMATION FOR SEQ ID NO: 253:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 239 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Cancerous prostate

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 48..238

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 35..225

id HSCOCCO21

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 15..49

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 94

region 1..35 id HSCOCC021

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 27..238

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 1..212

id T32119

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 36..238

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 1..203

id T35494

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 49..238

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 13..202 id HUMHG5097

(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 51238 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 1188 id AA027882 est (ix) FEATURE:	
(A) NAME/KEY: sig_peptide (B) LOCATION: 78137 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.6 seq AFVXACVLSLIST/IY	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 253:	
AAGAGTAGGG TGCTGTGGTC TGAGCTAGAG GGTGAAGCTG GCGGASAGGA GGATGGGCGA	60
GCAGTCTGAA TGCCAGA ATG GRT AAC CGT TTT GCT ACA GCA TTT GTA ATD Met Xaa Asn Arg Phe Ala Thr Ala Phe Val Xaa -20 -15 -10	110
GCT TGT GTG CTT AGC CTC ATT TCC ACC ATC TAC ATG GCA GCC TCC ATT Ala Cys Val Leu Ser Leu Ile Ser Thr Ile Tyr Met Ala Ala Ser Ile -5 1 5	158
GGC ACA GAC TTC TGG TAT GAA TAT CGA AGT CCA GTT CAA GAA AAT TCC Gly Thr Asp Phe Trp Tyr Glu Tyr Arg Ser Pro Val Gln Glu Asn Ser 10 20	206
AGT GAT TTG AAT AAA AGC ATC TGG GAT GAA TTG Ser Asp Leu Asn Lys Ser Ile Trp Asp Glu Leu 25 30	239
(2) INFORMATION FOR SEQ ID NO: 254: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 477 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR	
(ii) MOLECULE TYPE: CDNA	
(vi) ORIGINAL SOURCE:(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate	
<pre>(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(43130) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97</pre>	

est

(ix)	FEATURE:
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(A) NAME/KEY: other

(B) LOCATION: complement(137..219)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100 region 88..170 id C01485

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 421..459

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.6

seq MSLTSGFLRVSQG/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 254:

CACCAATGTT ATGAATGGCG TGGCCTCCTA CTGCCGTCCC TGTGCCCTAG AAGCCTCTGA 60 TGTGGGCTCC TCCTGCACCT CTTGTCCTGC TGGTTACTAT ATTGACCGAG ATTCAGGAAC 120 CTGCCAMTCC BTGCCCCCCT AACACAATTC TGAAAGCCCA CCAGCCTTAT GGTGTCCAGG 180 CCTGTGTGCC CTGTGGTCCA GGGACCAAGA ACAACAAGAT CCACTCTCTG TGCTACAATG 240 ATTGCACCTT CTCACGCAAC ACTCCAACCA GGACTTTCAA CTACAACTTC TCCGCTTTGG CAAACACCGT CACTCTTGCT GGAGGGCCAA GCTTCACTTC CAAAGGGTTG AAATACTTCC ATCACTTTAC CCTCAGTCTC TGTGGAAACC AGGGTAGGAA AATGTCTGTG TGCACCGACA 420 ATG TCA CTG ACC TCC GGA TTC CTG AGG GTG AGT CAG GGT TCT CCA AAT 468 Met Ser Leu Thr Ser Gly Phe Leu Arg Val Ser Gln Gly Ser Pro Asn CTA TCA CAG 477 Leu Ser Gln 5 -

(2) INFORMATION FOR SEQ ID NO: 255:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 315 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other

(B) LOCATION: 55..316

(C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 99 region 1..262 id H87671 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 102..261 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 50..209 id N47067 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 55..104 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 2..51 id N47067 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 251..316 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 14..79 id AA135001 est (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 58..246 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.6 seq AIRTLFSVTGILA/EO (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 255: AACTTGGCGC GCGGCSSGGC TGCAGACGGC TGCGAGGCGC TGGGCACAGG TGTCCTG 57 ATG GCA AAT TTC AAG GGC CAC GCG CTT CCA GGG AGT TTC TTC CTG ATC Met Ala Asn Phe Lys Gly His Ala Leu Pro Gly Ser Phe Phe Leu Ile ATT GGG CTG TGT TGG TCA GTG AAG TAC CCG CTG AAG TAC TTT AGC CAC Ile Gly Leu Cys Trp Ser Val Lys Tyr Pro Leu Lys Tyr Phe Ser His 153 ACG CGG AAG AAC AGC CCA CTA CAT TAC TAT CAG CGT CTC GAG ATC GTC Thr Arg Lys Asn Ser Pro Leu His Tyr Tyr Gln Arg Leu Glu Ile Val 201 -25 GAL GCC GCA ATT AGG ACT TTG TTT TCC GTC ACT GGG ATC CTG GCA GAG Glu Ala Ala Ile Arg Thr Leu Phe Ser Val Thr Gly Ile Leu Ala Glu 249 -15 -10 -5

CAG TTT GTT CCG GAT GGG CCC CAC CTG CAC CTC TAC CAT GAG AAC CAC
Gln Phe Val Pro Asp Gly Pro His Leu His Leu Tyr His Glu Asn His

5
10
15

TGG ATA AAG TTA ATG AAT Trp Ile Lys Leu Met Asn 20

315

(2) INFORMATION FOR SEQ ID NO: 256:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 405 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 89..405
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 84..400 id N34255

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 5..88
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 1..84 id N34255

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 89..304
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 83..298 id H79944

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 8..54
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 93 region 2..48

id H79944

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 336..382

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 91

region 332..378 id H79944

est

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 304..340

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 299..335 id H79944

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 54..88

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 49..83

id H79944

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 109..298

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 106..295

id H73369

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 2..88

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 1..87

id H73369

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 336..382

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 91

region 336..382

id H73369

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 295..326

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 293..324

id H73369

est

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(ix) FEATURE:
```

(A) NAME/KEY: other

(B) LOCATION: 164..237

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 142..215

id AA132425

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 327..395

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 307..375

id AA132425

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 21..88

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 94

region 3..70 id AA132425

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 124..163

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 92

region 103..142

id AA132425

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 109..298

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 24..213

id R97376

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 296..405

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 212..321

id R97376

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 187..342

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.5

WO 99/06550 PCT/IB98/01232

seq AGLLFGSLAGLGA/YO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 256:

AGCAGG	CACA A	ACAG	AGCC	GC T	cccc	WCTC	с тс	GCCC	CGCC	ACC	GGGA	CGG	AGAG	CGCCC	G 60
CCGCTG	CATT	TCCG	GCGA	CA C	CTCG	CAGG	т СА	TTCC	TGCG	GCT	TGCG	CGC	CCTT	GTAGA	120
AGCCGGGGCC TTCGTSAGAC CGGTGCAGGC CTGGGGTAGT CTCCTGTCTG GACAGAGAAG 18												180			
AGAAAA	ATG (Gln.	GAC Asp -50	ACT Thr	GGC Gly	TCA Ser	Val	GTG Val -45	CCT Pro	TTG (Leu	CAT His	Trp	TTT (Phe (-40	GGC Gly	228
TTT GGG	TAC Tyr	GCA Ala -35	GCA Ala	CTG Leu	GTT Val	GCT Ala	TCT Ser -30	GGT Gly	GGG Gly	ATC Ile	ATT Ile	GGC Gly -25	TAT Tyr	GTA Val	276
AAA GCA Lys Ala	GGC Gly -20	AGC Ser	GTG Val	CCG Pro	TCC Ser	CTG Leu -15	GCT Ala	GCA Ala	GGG G1y	CTG Leu	CTC Leu -10	TTT Phe	GGC Gly	AGT Ser	324
CTA GCC Leu Ala	GIA	CTG Leu	GGT Gly	GCT Ala	TAC Tyr 1	CAG Gln	CTG Leu	TCT Ser	CAG Gln 5	GAT Asp	CCA Pro	AGG Arg	AAC Asn	GTT Val 10	372
TGG GTT Trp Val	TTC Phe	CTA Leu	GCT Ala 15	ACA Thr	TCT Ser	GGT Gly	ACC Thr	TTG Leu 20	GCT Ala						405

(2) INFORMATION FOR SEQ ID NO: 257:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 323 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 119..237
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 94 region 116..234

id HSC2TH021

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 25..95
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 94

region 24..94 id HSC2TH021

est

(ix) FEATURE:

- (A) NAME/KEY: other (B) LOCATION: 238..289
- (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 90

region 234..285 id HSC2TH021

est

(ix) FEATURE:

- (A) NAME/KEY: other (B) LOCATION: 280..319
- (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 92 region 277..316 id HSC2TH021

est

(ix) FEATURE:

- (A) NAME/KEY: other (B) LOCATION: 130..237
- (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 23..130

id R59681 est

(ix) FEATURE:

- (A) NAME/KEY: other (B) LOCATION: 238..289
- (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 90 region 130..181 id R59681

est

(ix) FEATURE:

- (A) NAME/KEY: other (B) LOCATION: 280..325
- (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 93

region 173..218 id R59681

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide (B) LOCATION: 183..287
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 5.4

seq CCALLTSLXCIWG/PA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 257:

TCC	CGM	TCC	TTAT	GCTG	AT T	АТАА	CAAA	т сс	CTGD	RCCG	AAG	STAC	ттт	GATG	CTGCCG	120
GGA	RGCI	'GAC	TCCT	'GAGT	TC T	CACA	ACGC	T TG	ACCA	АТАА	GAT	TCGG	GAG	CTTC	TTCAGC	180
	ATG Met -35	GAG Glu	AKA Xaa	GGC (Leu :	AAA ' Lys : -30	TCA (Ser /	GCA Ala	GAC (Pro 1	CGG (Arg /	GAT (GGC Gly	ACC (GGT Gly	227
TAC Tyr ~20	ACT Thr	GRC Xaa	TTN Xaa	NKC Xaa	ARG Xaa -15	TAT Tyr	TGC Cys	TGT Cys	GCT Ala	TTA Leu -10	CT T Leu	ACA Thr	TCT Ser	TTA Leu	TGR Xaa -5	275
TGT Cys	ATT	TGG Trp	GGA Gly	CCT Pro 1	GCC Ala	TAC Tyr	CTA Leu	CAG Gln 5	TTA Leu	GCA Ala	CAT His	GGC Gly	TAT Tyr 10	GTA Val	AAG Lys	323

(2) INFORMATION FOR SEQ ID NO: 258:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 240 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 1..241

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 12..252

id H64050

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 1..241
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 1..241

id R17172

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 1..241
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 2..242

id HSC15C081

- (ix) FEATURE:
 - (A) NAME/KEY: other

WO 99/06550		263	PCT/IB
(C)	LOCATION: 8241 IDENTIFICATION MEG OTHER INFORMATION		
(ix) FEAT	URE:		
(B) (C)	NAME/KEY: other LOCATION: 29241 IDENTIFICATION ME OTHER INFORMATION:		
(ix) FEAT	URE:		
• •	NAME/KEY: sig_pept	ide	
	LOCATION: 10135		
	IDENTIFICATION MET OTHER INFORMATION:	CHOD: Von Heijne mat	rix
(6)	OTHER INFORMATION	score 5.4 seq ITGVILLAVGIWG	:/kv
		ood 110.1551.011.0	77 114
(xi) SEQU	ENCE DESCRIPTION: S	SEQ ID NO: 258:	
GGGCTAGTC ATG G	CG TCC CCG TCT CGG la Ser Pro Ser Arg -40	AGA CTG CAG ACT AAA Arg Leu Gln Thr Lys -35	A CCA GTC ATT 51 FPro Val Ile -30
ACT TGT TTC AAG	AGC GTT CTG CTA A	C TAC ACT TTT ATT T	TC TGG ATC 99
Thr Cys Phe Lys	Ser Val Leu Leu I	le Tyr Thr Phe Ile F	The Trp Ile
ACT GGC GTT ATC	CTT CTT GCA GTT G	GC ATT TGG GGC AAG G	STG AGC CTG 147
Thr Gly Val Ile -10	Leu Leu Ala Val G	ly Ile Trp Gly Lys V 1	al Ser Leu
GAG AAT TAC TTT	TCT CTT TTA AAT G	AG AAG GCC ACC AAT G	STC CCC TTC 195
Glu Asn Tyr Phe 5	Ser Leu Leu Asn G. 10	lu Lys Ala Thr Asn V	Val Pro Phe
GTG CTC ATT GCT	ACT GGT ACC GTC A	TT ATT CTT TTG GGC A	ACC TTG 240
Val Leu Ile Ala	Thr Gly Thr Val I.	le Ile Leu Leu Gly T	Thr Leu

(2) INFORMATION FOR SEQ ID NO: 259:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 385 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 8..349
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 6..347 id AA075824

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 344..385
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 341..382 id AA075824

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 22..366
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 2..346

id R55598

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 71..385
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 1..315 id HSC33B061

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 156..385
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 91..320

id T65515

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 70..141
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 5..76

id T65515

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 29..305
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99 region 2..278

id HSCZRF061 est

1	[ix]	FEATURE	•
٦	^	r Du I O IV D	٠

(A) NAME/KEY: sig_peptide

(B) LOCATION: 119..319

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.2

seq LSVSLLPCAGAWS/LL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 259:

AAAAGCGGAG	MYAGGMNGGG TGAG	GAGAGT CGAGGGAGG	T GACGCGCGCT GCCGGGGCGA	60
GGTTGCGAGG	GGCGGTGTTG AAGA	ATGTGT GGGCGAACA	T CCTGTCACTT ACCTAGAG	118
			C GGC GGG GCC AGC e Gly Gly Ala Ser Ser -55	166
		r Leu Ile Ser Th	I GCC CCT CTT ACA ACA r Ala Pro Leu Thr Thr -40	214
			T CTC GAC TGC ACT TTG s Leu Asp Cys Thr Leu -20	262
			G CTC CCA TGC GCT GGG u Leu Pro Cys Ala Gly -5	310
			C CTT CCA AGC TGC AGT e Leu Pro Ser Cys Ser 10	358
	MCA ACT GTC GT Xaa Thr Val Va 2	l Val Ile		385

(2) INFORMATION FOR SEQ ID NO: 260:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 386 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 43..128
 - (C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97 region 19..104 id R49759 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 132..194 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 106..168 id R49759 est (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 225..311 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5.1 seq LLMLGVTLPNSYW/RV (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 260: ATTCCTCTGA CCTGCCAGGA AGCAGAGAGA CCCACAGAGC AGGCAGGGAG GCAGAAAGTG 60 GAGACGGACC TGAGCCCGAG GAAGAGGCAG GCAGAGGCTG AGGCTGATTC CACCCCAGCC 120 TGCCTGGRAC AAACCCTCCT TAGCCGCAGC CCCTTCCAGT TCCCTAGGGG TTCTGCCCCT 180 CCCCCTCTCT GGGGCACCAG CCCCCCAGGG TCCTGCATCC NACC ATG TCG ATG GCT 236 Met Ser Met Ala GTG GAA ACC TTT GGC TTC TTC ATG GCA ACT GTG GGG CTG CTG ATG CTG 284 Val Glu Thr Phe Gly Phe Phe Met Ala Thr Val Gly Leu Leu Met Leu -20 GGG GTG ACT CTG CCA AAC AGC TAC TGG CGA GTG TCC ACT GTG CAC GGG 332 Gly Val Thr Leu Pro Asn Ser Tyr Trp Arg Val Ser Thr Val His Gly AAC GTC ATC AHC ACC AAC AHC ATC TTC GAG AAC CTC TGG TTT AGC AGT 380 Asn Val Ile Xaa Thr Asn Xaa Ile Phe Glu Asn Leu Trp Phe Ser Ser 15 GCC GGG 386 Ala Gly 25 (2) INFORMATION FOR SEQ ID NO: 261: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 222 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

	(VI) ((A)	ORGA TISS	ANIS	4: H				ic p	rosta	ate				
	(:	ix)	(B) (C)	URE: NAME LOCA IDEN OTHE	ATION NTIF:	N: 1: [CAT]	81 1 NO	метно	ide: reg:	olast ntity ion 1	7 94 112.	. 216				
	(;)	ix) I	(B) (C)	JRE: NAME LOCA IDEN OTHE	TION TIF	N: 13	18 ON 1	ЧЕТНО	ider regi	olast ntity ion l NAO55	, 97 120	. 158				
	(1	lx) E	(B) (C)	JRE: NAME LOCA IDEN OTHE	TION TIFI	N: 55	011 ON N	L4 1ETHO	D: V							
	(>	(i) S	SEQUE	ENCE	DESC	CRIP	CION	: SE(Q ID	NO:	261	:				
ACT(CAGAA	AGC 1	rtgg <i>i</i>	ACCGO	CA TO	CCTA	GCCG	C CG/	ACTC	ACAC	AAG	GCAG <i>l</i>	ABT 1	rgcc	ATG Met -20	57
				GTG Val -15												105
AKC Xaa	TGG Trp	CCA Pro	SSG Xaa 1	GAT Asp	ACC Thr	ACA Thr	GTC Val 5	AAA Lys	CCT Pro	GGA Gly	GCC Ala	AMA Xaa 10	AAG Lys	GAC Asp	ACA Thr	153
AAG Lys	GAC Asp 15	TCT Ser	CGA Arg	SCC Xaa	AAA Lys	CTG Leu 20	CCC Pro	CAG Gln	ACC Thr	CTC Leu	TCC Ser 25	AGA Arg	GGT Gly	TGG Trp	GGT Gly	201
				TGG Trp												222
(2)	INFO	ORMA1	rion	FOR	SEQ	ID I	NO: 1	262:								
	(i	i) S	(A) (B) (C)	CE C LENG TYPE STRA TOPO	STH: S: NU ANDE	366 JCLEI ONESS	base C AC S: DC	e pai CID DUBLE		,						

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 207..326
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 85..204 id W69716

10 W69/16

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 122..208
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 1..87 id W69716

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 316..366
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 195..245

id W69716

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 282..366
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 94..178

id W73842

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 207..287
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 17..97

id W73842

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 257..326
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 42..111

id W58108

(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 317366 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 90 region 101150 id W58108 est																
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 112312 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 5</pre>																
	(>	(i) S	EQUE	NCE	DESC	CRIPT	CION:	SEC) ID	NO:	262:					
ATA	AGGCC	CTC F	AGGGT	CCTC	er ti	TCCC	CTGGC	СТС	TTC	AGA	GGGG	CCG	rgg A	MCAC	GTCGC	60
AGTO	CGT	GCT 1	TTTAT	rgga <i>i</i>	AA CO	CAGGT	rg t g1	GAC	SCCGA	ATG	CCTO	GCCA	GC (CAC His	117
							CTG Leu									165
							CAC His									213
							TGC Cys									261
							GAG Glu -10									309
							AGC Ser									357

366

(2) INFORMATION FOR SEQ ID NO: 263:

TAT GCA TTG

Tyr Ala Leu

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 316 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Hypertrophic prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 121..264
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 127..270

id N24991

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 3..124
- (C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 10..131

id N24991

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 161..292
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 15..146

id HSC1WG111

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 176..310
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 1..135

id AA001396

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 176..265
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 1..90

id AA017578

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 191..265
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 1..75

id R17530

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 167..295
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4.9

seq GLWLALVDGLVRX/AP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 263:

ACTITITCCT ACGCAGCCGC TCCTGCCGCC GTGGTCGCTG GAGCTTTGCC TCTCTAGGCC	60
GGCAGCGCCT CTCCTCCATG GTCCTGTCTG TCAGCGCTGT TTTGGGAGCC CGCCGGTGAG	120
GCCGGGCCAC GCTCAGACAC TTCGATCGTC GAGTCTGTCA CTGGGC ATG GCG GGT Met Ala Gly	175
CAG TTC CGC AGC TAC GTG TGG GAC CCG CTG CTG ATC CTG TCG CAG ATC Gln Phe Arg Ser Tyr Val Trp Asp Pro Leu Leu Ile Leu Ser Gln Ile -35 -30 -25	223
GTC CTC ATG CAG ACC GTG TAT TAC GGC TCG CTG GGC CTG TGG CTG GCG Val Leu Met Gln Thr Val Tyr Tyr Gly Ser Leu Gly Leu Trp Leu Ala -20 -15 -10	271
CTG GTG GAC GGG CTA GTG CGA ASA GCC CCT CGC TGG ATC SCA GGG Leu Val Asp Gly Leu Val Arg Xaa Ala Pro Arg Trp Ile Xaa Gly	316

(2) INFORMATION FOR SEQ ID NO: 264:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 331 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 72..312
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 76..316

id W03477

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 2..78
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 7..83

id W03477 est

•

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 72..328

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 69..325 id W40364

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 3..78

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 1..76 id W40364

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 164..328

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 172..336

id R71313

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 72..158

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 80..166

id R71313

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 7..78

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 94

region 16..87

id R71313

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 164..328

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 151..315

id H87810

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 72..158

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 59..145

id H87810

est

(ix) FEATURE:

(B) (C)	NAME/KEY: other LOCATION: 1478 IDENTIFICATION METHOOTHER INFORMATION:	DD: blastn identity 98 region 266 id H87810 est											
(B) (C)	NAME/KEY: other LOCATION: 72274 IDENTIFICATION METHO	AME/KEY: other OCATION: 72274 DENTIFICATION METHOD: blastn											
(B) (C)	URE: NAME/KEY: other LOCATION: 2078 IDENTIFICATION METHO OTHER INFORMATION:	DD: blastn identity 96 region 159 id AA135694 est											
(B) (C)	URE: NAME/KEY: other LOCATION: 270328 IDENTIFICATION METHO OTHER INFORMATION:	DD: blastn identity 96 region 249307 id AA135694 est											
(B) (C) (D)	URE: NAME/KEY: sig_peptic LOCATION: 62295 IDENTIFICATION METHO OTHER INFORMATION: ENCE DESCRIPTION: SEC	DD: Von Heijne matrix score 4.9 seq VGAVFGLTTCISA/HV											
AGGCTGCCCT TGCG	CTTCCC GAGCTGGCGG GG	CCGTGGT GCGGGATCGA GATT	rgcgggc 60										
Met Ala Pro I	ys Val Phe Arg Gln T	AC TGG GAT ATC CCC GAT (yr Trp Asp Ile Pro Asp (70 -65	GGC ACC 109 Gly Thr										
GAT TGC CAC CGC Asp Cys His Arg -60	: AAA GCC TAC AGC ACC 1 Lys Ala Tyr Ser Thr -55	ACC AGT ATT GCC AGC GTG Thr Ser Ile Ala Ser Val -50	C GCT 157 l Ala										
GGC CTG ACC GCC Gly Leu Thr Ala -45	GCT GCC TAC AGA GTC Ala Ala Tyr Arg Val -40	ACA CTC AAT CCT CCG GGG Thr Leu Asn Pro Pro Gly -35	C ACC 205 y Thr										
TTC CTT GAA GGA Phe Leu Glu Gly	GTG GCT AAG GTT GGA Val Ala Lys Val Gly	CAA TAC ACG TTC ACT GCAGIN Tyr Thr Phe Thr Ala	A GCT 253 a Ala										

WO 99/06550 PCT/IB98/01232

-20

301

-25

GCT GTC GGG GCC GTG TTT GGC CTC ACC ACC TGC ATC AGC GCC CAT GTC

Ala Val Gly Ala Val Phe Gly Leu Thr Thr Cys Ile Ser Ala His Val
-10 -5 1

CGC GAG AAG CCC GAC GAC CCC CTG AAC CGG
Arg Glu Lys Pro Asp Asp Pro Leu Asn Arg
5
10

(2) INFORMATION FOR SEQ ID NO: 265:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 215 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Cancerous prostate

(ix) FEATURE:

-30

- (A) NAME/KEY: other
- (B) LOCATION: complement (44..183)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 1..140 id N78549

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(2..34)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 93

region 150..182

id N78549

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement (103..214)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 95

region 100..211

id N27605

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 150..203
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4.9

seq WLQVLPVILLLLG/VP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 265:

AGAGAGAGG GCCGCTACGC CGCACAGCAA ACAAGCTCCG CGACGTTTCC AGGACCCGGA 60 TAATCCCGCC CTTAGAGCAG AGCCGGAAGA AGGCGGGACG AACCGGAAGA GGGTGAAATG CTTTCGGTAG GCACTCCACG GCTGTGAAG ATG GCG GCG GCT GCG TGG CTT CAG 173 Met Ala Ala Ala Trp Leu Gln -15GTG TTG CCT GTC ATT CTT CTG CTT CTG GGA GTC CCC CCG TCG 215 Val Leu Pro Val Ile Leu Leu Leu Gly Val Pro Pro Ser -5 (2) INFORMATION FOR SEQ ID NO: 266: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 127 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Cancerous prostate (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(1..124) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 59..182 id AA045287 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement(1..124) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 45..168 id R77973 est (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (1..124) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 70..193 id AA136043 est (ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement(1..124)(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 60..183 id AA115201

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(1..124)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 60..183

id R72616

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: $5..\overline{1}15$
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4.8

seq LLILDMNVLYTDA/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 266:

ATAG ATG GAA ATA TAC TTT ATA TTT TGT ATC ATC GTG CCT ATA GCC GCT

Met Glu Ile Tyr Phe Ile Phe Cys Ile Ile Val Pro Ile Ala Ala

-35

-30

-25

GCC ACC GTG TAT AAA TCC TGG TGT CTG CTC CTT ATC CTG GAC ATG AAT
Ala Thr Val Tyr Lys Ser Trp Cys Leu Leu Leu Ile Leu Asp Met Asn
-20 -15 -10

GTA TTG TAC ACT GAC GCG TCC CCA CTC GGG Val Leu Tyr Thr Asp Ala Ser Pro Leu Gly

127

(2) INFORMATION FOR SEQ ID NO: 267:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 220 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (3) LOCATION: 48..140
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 91 region 36..128 id AA054941

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 130..197

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 92

region 117..184 id AA054941

est

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 48..218

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 94

region 36..206

id W68324

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 48..141

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 91

region 22..115

id H72703

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 130..218

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93

region 103..191

id H72703

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 29..59

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93

region 4..34

id H72703

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 48..140

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 91

region 16..108

id AA128297

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 130..218

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93 region 97..185

id AA128297

est

	(i)	(B)	TURE: NAM LOC TDE OTH	E/KE ATION	N: 4 CAT	81 ION	41 METH	ide reg	ntit	y 91 13					
	(ix	(B)	TURE: NAM LOC IDE OTH	ATION NTIFI	V: 1	30 ION 1	METH	ide reg	ntit	y 93 94:	182				
	(ix	(B) (C)	TURE: NAMI LOCA I DEI OTHI	ATION NTIFI	: 7: CAT:	ll	63 METH(DD: N	ce 4	. 8	ne ma				
	(xi) SEQU	JENCE	DESC	RIP	rion	: SE	QĮID	NO:	267	:				
ACT	GTCGAC	G TGTT	CTTC	CG GI	GGC	GGAC	G GC	GGAT'	ragc	CTT	CGCG	GGG (CAAA	ATTGRA	60
RCY(CDRGGC	C ATG Met	AGC Ser 2	AGA I Arg I	AT I	ACC A	Ser 1	CCA (Pro \ -25	GTG A	AAC (Asn 1	CCA (Pro 1	Ala '	GTC : Val :	ľTC ?he	109
CCC Pro	CAT C	TG ACC eu Thr -15	Val	GTG Val	CTT Leu	TTG Leu	GCC Ala -10	ATT Ile	GGC Gly	ATG Met	TTC Phe	TTC Phe -5	ACC Thr	GCC Ala	157
IGG Irp	TTC T Phe P	TC GT1 he Val 1	TAC Tyr	GAG Glu	GTC Val 5	ACC Thr	TCT Ser	ACC Thr	AAG Lys	TAC Tyr 10	ACT Thr	CGT Arg	GAT Asp	ATC Ile	205
_	AAA G Lys G														220
(2)	INFOR	MATION	FOR	SEQ	ID 1	NO: 2	268:								
	(i)	(B) (C)	NCE (LENC TYPE STRA	STH: E: NU ANDED	422 CLEI NESS	base C AC C DC	pai ID UBLE								
	(ii) MOLE	CULE	TYPE	: CI	ONA									

(vi) ORIGINAL SOURCE:

WO 99/06550	279	PCT/I
(A) (F)	ORGANISM: Homo Sapiens TISSUE TYPE: Cancerous prostate	
(B) (C)	JRE: NAME/KEY: other LOCATION: 135179 IDENTIFICATION METHOD: blastn OTHER INFORMATION: identity 91 region 1559 id R68571 est	
(B) (C)	URE: NAME/KEY: sig_peptide LOCATION: 309413 IDENTIFICATION METHOD: Von Heijne matrix OTHER INFORMATION: score 4.8 seq LMLSSSLPLLIWL/KD	
(xi) SEQUE	ENCE DESCRIPTION: SEQ ID NO: 268:	
AACTTTAGCC TCTGA	ATTGCA GGCCACCACT TCATTTACAT GGGGTGAGCA CCAATGCGTT	60
TTGTTCAATT CTTT	STTCAA AACCCCAAGA ATCTGGACAA CTTGCACTCA AGACCCTCTA	120
CGGGTTTGGC GAGCO	CAGTCC TTCAGTGGCT GTTTTCTAGT AGCTCCTTGG CAATTGAGGG	180
GAACTGGCTG GGACG	CACTCT CCAGTGCTGT CTGAAGGCCA AGGAGTGAAC AGGGATGGCT	240
GCCCTGCCTT GAAG	AGGGAA GGACTCTTTT CTATCCTTTC CAGCTATAGT CCCTGATCCC	300
TACATGTG ATG CGG Met Arg -35	G TTG GCA GCG GAA GCT CAT CCT GGG CGA ACT CAC ACA g Leu Ala Ala Glu Ala His Pro Gly Arg Thr His Thr -30 -25	350
	CTT AAA CCT TTT CTT ATG CTA AGT TCT TCC CTT CCC Leu Lys Pro Phe Leu Met Leu Ser Ser Ser Leu Pro -15	398
	CTA AAG GAC AGA Leu Lys Asp Arg 1	422
(2) INFORMATION	FOR SEQ ID NO: 269:	
(A) (B) (C)	NCE CHARACTERISTICS: LENGTH: 261 base pairs TYPE: NUCLEIC ACID STRANDEDNESS: DOUBLE TOPOLOGY: LINEAR	

- (2) INFORMA
 - (i) SE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 2..261

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93 region 17..276

id N23506 est

е

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 2..220

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 92

region 8..226 id R74310

est

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 219..261

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 226..268

id R74310

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 103..261

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 125..283

id N42319

est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 103..261

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 119..277

id N33735

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 105..261

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 143..299

id R23867

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 97..213

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 4.8

seq IILFSAIVGFIYG/YV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 269:

AAGI	rgcci	RRA (CCTTA	AGCCC	CT CA	ACGGT	CCTI	AA C	STCTO	CGGT	CGCC	CTC	SCC :	rckc <i>i</i>	AGCCTG	60
CCVE	3CCG(CGC 1	rcrko	CTGSS	SC GA	ACTCO	CTCAG	S SCA	AGCC					CTG Leu -35		114
				CAG Gln												162
				ATT Ile												210
				GAA Glu												258
GGA Gly																261

(2) INFORMATION FOR SEQ ID NO: 270:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 353 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(154..354)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 70..270

id AA164185

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 28..111
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 298..381

id AA164184

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 297..344
 - (C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER	INFORMATION:	score 4.8	
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seq SKVLFCSFSNVLG/FD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 270:

CCAACGTGTG CTTTGAAAAA AAGAAGGGAT GTTTTCTGTG TCAAATGAAG GTAATCATAG	60
ATCAAATTTG CTTATTGTCT TGTTCAAATC CTAGAAAACC ATTAGCATTT TTCTTTGCTT	120
GTAATATKAG AATCTAACAC TCATACAGAA TATTGGAAAG GTTACCCTAC AATTGTAAAT	180
TTGAAATTCT CCTTCTAATT CTGTCAGTTA TTTATTGACA TAGTAGTGGT TCTGTAGTCA	240
ÄGTGCATATA AGGTTTTGAA TGTTACATCT TATTNNNGGA TTWTTATTTT ATCATT ATG Met	299
GAG TAT AGC AAA GTT CTA TTT TGT TCT TTT TCA AAT GTA CTT GGT TTT Glu Tyr Ser Lys Val Leu Phe Cys Ser Phe Ser Asn Val Leu Gly Phe -15 -5 1	347
GAT TAT Asp Tyr	353

(2) INFORMATION FOR SEQ ID NO: 271:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 225 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 19..133
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 1..115

id HSC13B041

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 124..226
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 105..207

id HSC13B041

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 124..226

20

(C) IDENTIFICATION METHOD: blastn
(D) OTHER INFORMATION: identity 96

region 71..173

id T08849 est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 53..133

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 1..81 id T08849

est

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 53..135

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 1..83 id H88132

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 124..192

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 92

region 71..139 id H88132

est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 192..226

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 94

region 140..174

id H88132

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 53..144

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 1..92 id T33149

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 145..226

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 92..173

id T33149

est

(ix) FEATURE:

			(B) (C)	LOC	E/KE ATIO NTIF ER I	N: 5 ICAT	21 ION	33 METH	ide reg	blas ntit ion AA12	y 98 18					
	(ix)	FEAT		E/KE	Y: o	ther									
			(B)	LOC	OITA	N: 1	92									
					NTIF: ER II					blas: ntit:						
			, - ,					J	reg.	ion	141.	.175				
									id i	AA12	1114					
	,		CC MO	unn												
	(:	1X)	FEAT		E/KE	Y: s:	ig pe	eptio	de							
			(B)	LOCA	OITA	N: 4	51	23								
			(C) (D)	OTHE	NTIFI ER IN	ICAT. NFORI	ION I ATIO	METHO ON:		Jon I ce 4.		ne ma	atri	ĸ		
			• •					••••		LIM(VLLT	RC/PI	:		
	(:	ki) :	SEQUI	ENCE	DES	CRIP	NOI	: SE	O ID	NO:	271	•				
												•				
ACT	CTCT	GAC	TGGG	GTGA	GG C	CGCA	GCGG	A CT	GCCC'	TTTC	CCA	AG A'	rg g	CG T	CG AAG	G 57
													et A		er Lys	
ATA	GGT	TCG	AGA	CGG	TGG	ATG	TTG	CAG	CTG	ATC	ATG	CAG	TTG	GGT	TCG	105
шe	GIY	-20	Arg	Arg	Trp	Met	Leu -15	Gln	Leu	Ile	Met	Gln -10	Leu	G1y	Ser	
cmc	O.M.O.	000					-									
Val	Leu	Leu	ACA Thr	CGC	TGC	Pro	TTT	TGG	GGC	TGC	TTC	AGC	CAG	CTC	ATG	153
	-5			,		1			01,	5	1.1.0	Jer	GIN	Leu	10	
CTG	TAC	GCT	GAG	AGG	GCT	GAG	GCA	CGC	CGG	AAG	CCC	GAC	ATC	CCA	GTG	201
Leu	Tyr	Ala	Glu	Arg	Ala	Glu	Ala	Arg	Arg	Lys	Pro	Asp	Ile	Pro	Val	201
		•		15					20					25		
			TAT													225
Pro	Tyr	Leu	Tyr 30	Phe	Asp	Ser	Gly									
(2)	INFO	ORMA'	TION	FOR	SEQ	ID N	10: 2	272:								
	(i) SE	EQUEN	ICE C	HARA	ACTEF	RISTI	CS:								
			(A)	LENG	TH: : NU	305	base	pai	rs							
			(C)	STRA	NDE	NESS	: DC	UBLE	Ξ							
					LOGY											
	(<u>i</u>	.i) P	MOLEC	CULE	TYPE	: CE	NA									
	()	/i) (ORIG1	NAL	SOUF	CE:										

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 2..287

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 9..294 id W52125

est

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 19..283

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 1..265 id AA024623 est

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 22..284

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 1..263 id H55824

est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 21..307

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 2..288 id R62921

est

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 102..287

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 57..242 id N31702

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 45..100

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 1..56 id N31702

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 69..224

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 4.7

seq LGLALGRLEGGSA/RH

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 272:

ATTGGCTCCG GA	TCGTGCGT GAGGCGGC	TT CGTGGGCAGC GA	GAGTCACA GACAAGACAG	60
CAAGCAGG ATG (GAG CAC TAC CGG A Glu His Tyr Arg L -50	AA GCT GGC TCT G ys Ala Gly Ser V -45	TA GAG CTC CCA GCG al Glu Leu Pro Ala ~40	110
CCT TCC CCA A: Pro Ser Pro Me	et Pro Gln Leu Pro	F CCT GAT ACC CT Pro Asp Thr Let -30	T GAG ATG CGG GTC u Glu Met Arg Val -25	158
CGA GAT GGC AC Arg Asp Gly Se -20	GC AAA ATT CGC AAG er Lys Ile Arg Ass -19	n Leu Leu Gly Leu	G GCT CTG GGT CGG u Ala Leu Gly Arg -10	206
TTG GAG GGC GC Leu Glu Gly Gl -5	GC AGT GCT CGG CAT ly Ser Ala Arg His 1	GTA GTG TTC TCA Val Val Phe Ser 5	A GGT TCT GGC AGG r Gly Ser Gly Arg 10	254
GCT GCA GGA AAA Ala Ala Gly Ly	AG GCT GTC AGC TGC ys Ala Val Ser Cys 15	GGCT GAG ATT GTG Ala Glu Ile Val 20	C AAG CGG CGG GTC l Lys Arg Arg Val 25	302
CCG Pro				305

(2) INFORMATION FOR SEQ ID NO: 273:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 322 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 113..324
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 2..213

id W26501

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 111..324
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 6..219

		207
		d W28013 st
(ix)	i	: blastn
(ix)	i	
(ix)	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 104181 (C) IDENTIFICATION METHOD (D) OTHER INFORMATION: s	: Von Heijne matrix
(xi)	SEQUENCE DESCRIPTION: SEQ	ID NO: 273:
rttgc	AAAGATGGCT GTAGGAATGG AGGA	GCCTGT ATTGCCGCTA ATGTGTG
CACAA	GGCTTCACTG GACCCAGCTG TGA	NACGACA TTG ATG AAT GCT CT Met Asn Ala Le

AGC	ATTT'	rgc A	\AAG#	ATGGC	or G	PAGGA	AATGO	3 AGC	SAGCO	CTGT	ATTO	3CCG(CTA A	ATGTO	STGTGC	60	
CTG	CCAC	CAA (GCTT	CACT	rg g <i>i</i>	ACCC	AGCTO	G TG∤	AAAC	GACA	TTG		AAT Asn -25			115	
					GTG Val											163	
					AGT Ser											211	
					AAT Asn											259	
					CCA Pro											307	
		ATT Ile 45														322	

(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 337 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR											
(ii)	MOLECULE TYPE: CDNA											
(vi)	ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Hypertrophic prostate											
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 94339 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 62307 id AA133635 est											
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 3297 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 95 region 166 id AA133635 est											
(ix)	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 191325 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.6 seq VLVYLVTAERVWS/DD											
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 274:											
ACTCCCAGGC	TGGGCCAGCA CACCCGGCAG GCTCTGTCCT GGAAACAGGC TTCAACGGGC	60										
TTCCCCGAAA	ACCTTCCCCG CTTCTGGRTA TGAAVWTKCA AGCTGCTTGC TGAGTCCTAT	120										
TGCCGGCTGC	TGGGAGCMAG GAGAGCCCTG AGGAGTAGTC ACTCAGTAGC AGCTGACGCG	180										
1	ATG AAC TGG AGT ATC TTT GAG GGA CTC CTG AGT GGG GTC Met Asn Trp Ser Ile Phe Glu Gly Leu Leu Ser Gly Val -45 -40 -35	229										
AAC AAG TAC Asn Lys Tyr -30	TCC ACA GCC TTT GGG CGC ATC TGG CTG TCT CTG GTC TTC Ser Thr Ala Phe Gly Arg Ile Trp Leu Ser Leu Val Phe -25	277										
ATC TTC CGC Ile Phe Arg -15	GTG CTG GTG TAC CTG GTG ACG GCC GAG CGT GTG TGG AGT Val Leu Val Tyr Leu Val Thr Ala Glu Arg Val Trp Ser	325										
GAT GAC CAC Asp Asp His 1		337										

(2) INFORMATION FOR SEQ ID NO: 275:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 287 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR 	
(ii) MOLECULE TYPE: CDNA	
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Prostate</pre>	·
<pre>(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 205287 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 96</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 129176 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.5</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 275:	
ACTGTCCCAT TCCTCCCCCT ACAACACACA CACCTTTCAG GCAGGGASGN GATGAGCTTC	60
CAGCCCCAAG AGTGGAGGCT GCCACATCCT AACATASGKA KCTATTGRRA AGGAAKSAGT	120
GTGTATCT ATG ATT ATA TCT CTG TTC ATC TAT ATA TTT TTK ACA TGT AGC Met Ile Ile Ser Leu Phe Ile Tyr Ile Phe Xaa Thr Cys Ser -15 -10 -5	170
AAC ACC TCT CCA TCT TAT CAA KGA ACT CAA CTC GGT CTG GGT CTC CCC Asn Thr Ser Pro Ser Tyr Gln Xaa Thr Gln Leu Gly Leu Gly Leu Pro 1 5 10	218
AGT GCC CAG TGG TGG CCT TTG ACA GGT AGG AGG ATG CAG TGC TGC AGG Ser Ala Gln Trp Trp Pro Leu Thr Gly Arg Arg Met Gln Cys Cys Arg 15 20 25 30	266
CTA TTT TGT TTT KTG TTA CAA Leu Phe Cys Phe Xaa Leu Gln 35	287
(2) INFORMATION FOR SEQ ID NO: 276:	
(i) SEQUENCE CHARACTERISTICS:	

- (A) LENGTH: 156 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

- (A) NAME/KEY: other(B) LOCATION: 1..156
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98 region 40..195 id AA227366

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 1..156
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98 region 4..159 id AA069390

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 9..152
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100 region 1..144 id AA248850 est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 18..95
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96 region 1..78 id AA248912

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 88..132
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 93 region 70..114

id AA248912

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 61..108
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4.4

seq LNSLSALAELAVG/SR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 276:

ATGGCTGTCA GAGGTGGGCG GCTTTGACCG AGAGGCTGCT GGAGCTCGTG TTTGGACGCG 60

ATG TTT CGT CTG AAC TCA CTT TCT GCT TTG GCA GAA CTG GCT GTG GGT

Met Phe Arg Leu Asn Ser Leu Ser Ala Leu Ala Glu Leu Ala Val Gly

-15

-10

-5

TCT CGA TGG TAC CAT GGA GGA TCA CAG CCC ATC CAG ATC CGG CGG AGA

Ser Arg Trp Tyr His Gly Gly Ser Gln Pro Ile Gln Ile Arg Arg Arg

1 5 10 156

- (2) INFORMATION FOR SEQ ID NO: 277:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 369 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 98..330
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 69..301

id R99696

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 29..98
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 1..70 id R99696

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 206..330
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 164..288

id W90165

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (3) LOCATION: 98..209

WO 99/0	6550 · 292	PCT/IB9
	(C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 55166 id W90165 est	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 4398 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 96 region 156 id W90165 est	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 98330 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 82314 id H91200 est	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 1698 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 95 region 183 id H91200 est	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 98249 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 62213 id R06513 est	
(ix)	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 238288 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.4 seq TLRTWLCCAGSWA/VE	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 277:	
ATACTTGC	AGCTARAACT AAATATTGCT GCTTGGGGAC CTCCTTCTAG CCT	TAAATTT 60

CAGCTCATCA CCTTCACCTG CCTTGGTCAT GGCTCTGSCT ATTCTCCTTG ATCCTTGCCA 120

TTTGCACCAG ACCTGGATTC CTAGCGTCTC VATCTGGAGT GCGGCTGGTG GGGGGCCTCC 180

ATG ACG GCT GGG ACA TTA AGG ACG TGG CTG TGT TGT GCC GGG AGC TGG 285

237

ACCGCTGTGA AGGGCGGGTG GAGGTGGAAC AGAAAGGCCA GTGGGGCACC GTGTGTG

Met Thr Ala Gly Thr Leu Arg Thr Trp Leu Cys Cys Ala Gly Ser Trp
-15

GCT GTG GAG CTG CCA GCG GAA CCC CTA GTG GTA TTT TGT AWG AGC ACC
Ala Val Glu Leu Pro Ala Glu Pro Leu Val Val Phe Cys Xaa Ser Thr
1 5 10 15

AGC AGA AAA AGA GCA AAA GGT CTC ATC CAA TCA GTC
Ser Arg Lys Arg Ala Lys Gly Leu Ile Gln Ser Val
20
25

(2) INFORMATION FOR SEQ ID NO: 278:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 188 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(2..99)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 99..196 id AA088690

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(87..187)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95 region 12..112

id AA088690

est

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 111..182
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.3

seq RLLVILCVSVKAG/ST

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 278:

ACTACAGCAT GGCCACGTGG AGGCAGCGGC AGGAGAAAAA GCAGCTGGGC TTCTTCTGAA 60

CCCAAGCCCT CTCGACTGCC CCTATCCCCT GGAVCCCCAA CATACCTACA ATG CTG
Met Leu

GGG AGG CCC TGC TTC CAC TCC CCT CAG AGG CTT TTG GTC ATC CTC TGC 164

Gly Arg Pro Cys Phe His Ser Pro Gln Arg Leu Leu Val Ile Leu Cys
-20 -15 -10

GTG TCA GTA AAA GCA GGC AGC ACG Val Ser Val Lys Ala Gly Ser Thr -5

188

- (2) INFORMATION FOR SEQ ID NO: 279:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 289 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: CDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 106..261
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 119..274 id AA280906

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 2..99
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 15..112 id AA280906

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 260..291
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96

region 272..303

id AA280906

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 140..291
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 224..375

id HUM406F04B

- (ix) FEATURE:
 - (A) NAME/KEY: other

```
(B) LOCATION: 12..112
```

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 1..101 id HUM406F04B

est

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 106..140

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 94..128 id HUM406F04B

est

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 132..261

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 124..253 id AA133362

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 5..92

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 1..88 id AA133362

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 260..291

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96 region 251..282

id AA133362

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 106..261

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 94..249 id N57260

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 10..92

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 1..83 id N57260

	296	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 260291 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 96 region 247278 id N57260 est	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 41234 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 95 region 42235 id W25567 est	
(ix)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 140 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 140 id W25567 est	
(ix)	FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 194277 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.2 seq LQFVLPVATQIQQ/EV	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 279:	
	GGAACGGTTG TAGGACGTGG CTCTTTATTC GTGAGTTTTC CATTTACCTC	60
	AGAGCTTCAG ACGCCCTATG GCGTCCGCCT CGACACCAAC CGGCGGCCTT	120
	CAAGCAAAGG TGGTCCTCGC GGAGGTGATC CAGGCGTTCT CCGCCCCGGA	180
GAATGCAGTG (CGC ATG GAC GAG GCT CGG GAT AAC GCC TGC AAC GAC ATG Met Asp Glu Ala Arg Asp Asn Ala Cys Asn Asp Met -25 -20	229
GGT AAG ATG Gly Lys Met -15	CTG CAA TTC GTG CTG CCC GTG GCC ACG CAG ATC CAG CAG Leu Gln Phe Val Leu Pro Val Ala Thr Gln Ile Gln Gln -10 -5	277
GAG GTT ATC Glu Val Ile 1		289

(2) INFORMATION FOR SEQ ID NO: 280:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 355 base pairs

297
(B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR
(ii) MOLECULE TYPE: CDNA
<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Normal prostate</pre>
<pre>(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 3869 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 96 region 132 id H56508 est</pre>
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 287349 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 4.2</pre>
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 280:
AAACCTCCGT GGCTAGTCTT GACGTGGCGG GTTGCTTTCC AAAATGGCGC GGGTGCTGAA 60
GGCTGCAGCC GCDBAATGCC GTAGGTGAAT ACCGGGCACC GCCGACCTTC GCCATGGGAC 120
AGGGAGCGTG GGAACGGCGG TCGGGGGCGG AGGAKGCCTC GGTGTGGCCA AAGCACCTTG 180
ATCTAATGTC CTCCCCGGG GGCGCGTTCC ACAGCAGCTG CTGTCACTTW KGGCAGAGGG 24
TGCCTTCCAG AAGCGCCACC GCTTAGTAGC GGGGATTGCB TTGTGC ATG AGT CCC Met Ser Pro -20
ATT TCE ATC CGA GAG CTG TGC GCC TTG GGC TCT GCA CCT TCC AGT ATG 11e Ser Ile Arg Glu Leu Cys Ala Leu Gly Ser Ala Pro Ser Ser Met -15 -5
TGG GCB GGA GAG Trp Ala Gly Glu 1
(2) INFORMATION FOR SEQ ID NO: 281:
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 258 base pairs (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 108..255

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97 region 2..149

id AA095592

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 18..105

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 219..306

id T70757

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 163..255

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 37..129

id H66541

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 163..255

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 37..129

id R92835

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 172..255

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 14..97

id H87601

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 52..90

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 4.2

seq MTDLLSASPWALT/IV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 281:

		TCA Ser						105
		TCC Ser					 	153
		TTA Leu					 	201
		TCT Ser					 	249
 CAY His								258

(2) INFORMATION FOR SEQ ID NO: 282:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 285 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 6..202
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 10..206

id AA074428

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 193..254
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 95

region 196..257

id AA074428

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 28..202
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99 region 1..175

300

id AA158941 est PCT/IB98/01232

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 193..285
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 165..257

id AA158941

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 37..202
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 1..166 id AA148039

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 193..254
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 156..217

id AA148039

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 250..285
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 214..249

id AA148039

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 74..280
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 1..207

id H72224

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 76..153
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4.2

seq LTCGPALVPRLWA/TC

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 282:

AAGAGGCTAG AAGCTGGATT CAGCGTGTCC GCGACCTCAC CTTTAGGTCC TGTGAGGGAC 60
GGCCCAGGTG GCAGG ATG TCC TGG TCT GGC CTT CTC CAT GGC CTC AAC ACG 111

Met Ser Trp Ser Gly Leu Leu His Gly Leu Asn Thr
-25 -20 -15

TCC CTA ACT TGT GGC CCA GCT CTG GTT CCC CGG CTC TGG GCT ACC TGC

Ser Leu Thr Cys Gly Pro Ala Leu Val Pro Arg Leu Trp Ala Thr Cys

Ser Leu Thr Cys Gly Pro Ala Leu Val Pro Arg Leu Trp Ala Thr Cys
-10 -5 l

TCC ATG GCT ACC CTG AAC CAG ATG CAC CGC CTG GGG CCC CCC AAG CGG
Ser Met Ala Thr Leu Asn Gln Met His Arg Leu Gly Pro Pro Lys Arg

5 10 15

CCG CCT CGG AAG CTG GGC CCC ACG GAA GGC CGG CCG CAG CTG AAG GGT
Pro Pro Arg Lys Leu Gly Pro Thr Glu Gly Arg Pro Gln Leu Lys Gly
20 25 30

GTG GTC CTG TGC ACG TTT ACC CGC AAC CGG
Val Val Leu Cys Thr Phe Thr Arg Asn Arg
35 40

(2) INFORMATION FOR SEQ ID NO: 283:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 225 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 18..223
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 93 region 1..206 id HSC3CC061

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 112..223
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 91

region 94..205

id H33976

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 49..93
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 91 region 1..45 id AA041823

- (A) NAME/KEY: other
- (B) LOCATION: 49..93
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 93

region 1..45 id AA003782

(ix) FEATURE:

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 25..93
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4.1

seq LEAFSQAISAIQA/LR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 283:

- AAKAGCTGCT GTGGCGGCGG CAAC ATG GCG GAC GTG ATA AAT GTC AGT GTG

 Met Ala Asp Val Ile Asn Val Ser Val

 -20

 -15
- AAC CTG GAG GCC TTT TCC CAG GCC ATT AGT GCC ATC CAG GCG CTG CGA
 Asn Leu Glu Ala Phe Ser Gln Ala Ile Ser Ala Ile Gln Ala Leu Arg
 -10 -5
- TCC AGC GTG AGC AGG GTG TTC GAC TGC CTG AAG GAT GGG ATG CGG AAC

 Ser Ser Val Ser Arg Val Phe Asp Cys Leu Lys Asp Gly Met Arg Asn

 5 10 15
- AAG GAG ACG CTG GAG GGC CGG GAG AAG GCC TTT ATT GCG CAC TTC CAG
 Lys Glu Thr Leu Glu Gly Arg Glu Lys Ala Phe Ile Ala His Phe Gln
 20 25 30
- GAC AAC TTA CAT TCG GTC AAC CGG GAC CCA
 Asp Asn Leu His Ser Val Asn Arg Asp Pro
 35 40 225

(2) INFORMATION FOR SEQ ID NO: 284:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 339 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement (210..340)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 172..302

id AA062591

(ix) FEATUR

(A) NAME/KEY: sig peptide

(B) LOCATION: 109..204

- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4.1

seq RLLSSLLLTMSNN/NP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 284:

AGACCCGATG GACCCCGGCG ACGCSCCATT TTGGAGTCTT CCCTAAGGAT CCTCTACCGG 60

CTTTTCGAGT CAGTGCTGCC GCCGCTGCCC GCGGCTTTGC AGAGCAGG ATG AAT GTG Met Asn Val -30

ATA GAC CAC GTG CGG GAC ATG GCG GCC GCG GGG CTG CAC TCC AAC GTG 165

Ile Asp His Val Arg Asp Met Ala Ala Ala Gly Leu His Ser Asn Val -25 -20 -15

CGG CTC CTC AGC AGC TTG TTA CTT ACA ATG AGT AAT AAC AAC CCT GAG Arg Leu Leu Ser Ser Leu Leu Leu Thr Met Ser Asn Asn Asn Pro Glu -10 -5 1

TTA TTC TCC CCA CCT CAG AAG TAC CAG CTT TTG GTG TAT CAT GCA GAT 261

TTA TTC TCC CCA CCT CAG AAG TAC CAG CTT TTG GTG TAT CAT GCA GAT

Leu Phe Ser Pro Pro Gln Lys Tyr Gln Leu Leu Val Tyr His Ala Asp

5 10 15

TCT CTC TTT CAT GAT AAG GAA TAT CGG AAT GCT GTG AGT AAG TAT ACC
Ser Leu Phe His Asp Lys Glu Tyr Arg Asn Ala Val Ser Lys Tyr Thr
20 25 30 35

ATG GCT TTA CAG CAG AAG AAA GCG CTA AGT

Met Ala Leu Gln Gln Lys Lys Ala Leu Ser

40 45

(2) INFORMATION FOR SEQ ID NO: 285:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 141 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(2..41)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 90

region 66..105 id AA085310

est

1	ix)	FEATURE:	

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 70..117
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4.1

seq ACLAWTAVRPSAC/CH

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 285:

AAAGTGAGTT TGCGAACGGA GCAGCTGCTG CAGCAGGGCC CATGGCGGAC ACCCAGTACA 60

TCCTGCCCA ATG ACA TCG GCG TGT CTA GCC TGG ACT GCC GTG AGG CCT TCC 111

Met Thr Ser Ala Cys Leu Ala Trp Thr Ala Val Arg Pro Ser

-15 -10 -5

GCC TGC TGT CAC CCA CAG AGC GCC AAC TGG
Ala Cys Cys His Pro Gln Ser Ala Asn Trp

1 5

(2) INFORMATION FOR SEQ ID NO: 286:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 290 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement (147..290)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 90 region 141..284 id W12393
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 249..289
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 1..41 id HSC2TF111

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide(B) LOCATION: 60..224

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 4

seq VFGMSSSSGASNS/AP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 286:

ATCTCAACTT GGACTTGCAA TCACAGAACA TTTACCACCA TGGAAGAGAA GGAAGTAGG 59 ATG AAT GGA AGT AGG ACT CTA ACG CAC AGC ATT AGT GAT GGA CAA CTT 107 Met Asn Gly Ser Arg Thr Leu Thr His Ser Ile Ser Asp Gly Gln Leu -50 CAA GGT GGC CAG TCC AAT AGT GAA CTA TTT CAG CAG GAG SSA CAG ACA Gln Gly Gly Gln Ser Asn Ser Glu Leu Phe Gln Gln Glu Xaa Gln Thr -30 GCA CCA GCT CAA GTT CCT CAA GGC TTT AAT GTT TTT GGA ATG TCC AGT 203 Ala Pro Ala Gln Val Pro Gln Gly Phe Asn Val Phe Gly Met Ser Ser -15 TCC TCT GGT GCT TCA AAT TCA GCA CCA CAT CTT GGA TTT CAC TTA GGC 251 Ser Ser Gly Ala Ser Asn Ser Ala Pro His Leu Gly Phe His Leu Gly AGC AAA GGA ACA TCT AGC CTT TCT CAA CAA ACT CCC GGG 290 Ser Lys Gly Thr Ser Ser Leu Ser Gln Gln Thr Pro Gly 15

(2) INFORMATION FOR SEQ ID NO: 287:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 326 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement (68..194)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 204..330

id N35493

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement (208..323)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 75..190 id N35493

est

è	(ix)	FEATURE:
Į	1 X	L'EATURE:

- (A) NAME/KEY: other
- (B) **EOCATION**: complement(2..79)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 318..395

id N35493

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 186..233
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4

seq FFLFLSFVLMYDG/LR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 287:

ATAAAAGAAG CAGCAAATAG AATTTCCCAC AAAGTAAGTT GACTCTAAAT CTTAAGTATT 60

ACCTAGTTTT TTAAAGGTTT GAATATAATA ATGCAGTATT TGCAGTATAA AAAGGAAGGA 120

ATTTGTAGAG AATCATTTTG GTGCTCAAGT CTCTTAGCAG TGCCTTATTG CCTCATAGCA 180

AGAAG ATG CTG GGG TTT TTT TTG TTT TTG TCC TTT GTA TTA ATG TAT GAT 230 Met Leu Gly Phe Phe Leu Phe Leu Ser Phe Val Leu Met Tyr Asp -15 -5

GGT TTG CGC CTT TTT GGC ATT CTT TCA ACA TGT CGT GTA CAT CAC ACC
Gly Leu Arg Leu Phe Gly Ile Leu Ser Thr Cys Arg Val His His Thr
1 5 10

ATG AAT CAG TTC CTA ATT GAT ATA TCT AGC TTT ACC TCC CGA GTT CGG
Met Asn Gln Phe Leu Ile Asp Ile Ser Ser Phe Thr Ser Arg Val Arg
20 25 30

(2) INFORMATION FOR SEQ ID NO: 288:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 383 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 219..380
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 46..207

id N95583 est

ix		נוח		

- (A) NAME/KEY: other
 (B) LOCATION: 219..335
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 46..162 id AA283710

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 336..380
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100 region 162..206 id AA283710

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 240..320
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4

seq SIKVLLQSALSLG/RS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 288:

AGTGGCTCTT CTGACCCAAG GCCCCGCCGT CCAGGTAGGG GGCTGTGGCC TCTAGGGATC 60 AGGGACTACT TACCTGCGAA TCCCGGTTGC CCGCCCGCCA RCACGTCCGK TYCCSTAARG 120 CARAMCGCCT KGGCTCCTGG CTGAACCGTC TTCTCAMCGT TTGSCGGAGT CTGAMCTCCC CACGCTTAGT CCACTAACGR AGCTATCCCT GCTCCTGMCC CACAGCTTCT AAGTGCCAG 239 ATG ATG GAG GAG CGT GCC AAC CTG ATG CAC ATG ATG AAA CTC AGC ATC 287 Met Met Glu Glu Arg Ala Asn Leu Met His Met Met Lys Leu Ser Ile AAG GTG TTG CTC CAG TCG GCT CTG AGC CTG GGC CGC AGC CTG GAT GCG 335 Lys Val Leu Leu Gln Ser Ala Leu Ser Leu Gly Arg Ser Leu Asp Ala -5 GAC CAT GCC CCC TTG CAG CAG TTC TTT GTA GTG ATG GAG CAC TGC TCA 383 Asp His Ala Pro Leu Gln Gln Phe Phe Val Val Met Glu His Cys Ser

15

(2) INFORMATION FOR SEQ ID NO: 289:

10

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 319 base pairs
- (B) TYPE: NUCLEIC ACID
- (C) STRANDEDNESS: DOUBLE
- (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 57..180
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97 region 15..138

id AA090170

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 226..286
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 184..244 id AA090170

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 21..242
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 90

region 1..222 id HSU46267

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 143..319
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 90

region 220..396

id AA048294

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 149..286
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 92

region 139..276

id AA118611

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 143..286
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 92

region 88..231

id AA063937

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide
(B) LOCATION: 80..130

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 3.9

seq XIVSAALLAFVQT/HL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 289:

AGTTGGTGGG GCTGGGGGAT GAGAGCTGCA CCGCGCGGGA YAAGTCGCCG GCGGCGCCCG 60

AMGGAGCAGA ACAGAGAGC ATG GAG CTG GAG AKG ATC GTC AGT GCA GCC CTC 112

Met Glu Leu Glu Xaa Ile Val Ser Ala Ala Leu

-15

CTT GCC TTT GTC CAG ACA CAC CTC CCG GAG GCC GAC CTC AGT GGC TTG
Leu Ala Phe Val Gln Thr His Leu Pro Glu Ala Asp Leu Ser Gly Leu
-5 1 5 10

GAT GAG GTC ATC TTC TCC TAT GTG CKT GGG GTC CTG GAG GAC CTG GGC
Asp Glu Val Ile Phe Ser Tyr Val Xaa Gly Val Leu Glu Asp Leu Gly
15 20 25

CCC TCG GGC CCA TCA GAG GAG AAC TTC GAT ATG GAG GCT TTC ACT GAG
Pro Ser Gly Pro Ser Glu Glu Asn Phe Asp Met Glu Ala Phe Thr Glu
30 35

ATG ATG GAG GCC TAK GTG CCT GGC TTC GCC CAC ATC CCC AGG GGM ACA

Met Met Glu Ala Xaa Val Pro Gly Phe Ala His Ile Pro Arg Gly Thr

45 50 55

ATA GGG GAS ATG ATG

Ile Gly Xaa Met Met

60

(2) INFORMATION FOR SEQ ID NO: 290:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 274 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 2..273
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 93 region 8..279 id T30552

est

(ix) FEATURE:

WO 99/06550 PCT/IB98/01232

 (A) NAME/KEY: other (B) LOCATION: 3273 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 92 region 1271 id C14403 est 	
<pre>(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 2273 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 92</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 4273 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 92</pre>	
(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 3270 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 93 region 1268 id C14440 est	
(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 98175 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.9 seq SLIPLFXFIGTGA/TG	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 290:	
GTTCTCTCCK KTCCAGTCGG AGACCTCTGC SGVPPPC ATC CTG GGG GAG ATC	.15
GGT CAG GCC AAG AAG CAT CCG AGC TTG ATC CCC CTC TTT KTA TTT ATT Sly Gln Ala Lys Lys His Pro Ser Leu Ile Pro Leu Phe Xaa Phe Ile -20 -15 -10 -5	.63
GGR ACT GGA GCT ACT GGA GCA ACA CTG TAT CTC TTG CGT CTG GCA TTG 2 Sly Thr Gly Ala Thr Gly Ala Thr Leu Tyr Leu Leu Arg Leu Ala Leu 1 5 10	11
Phe Asn Pro Xaa Val Cys Trp Asp Arg Xaa Asn Pro Glu Pro Trp Asn	59

WO 99/06550 PCT/IB98/01232

				311			
		15	20		25		
•		GC CCC ly Pro				•	274
(2)	INFOR	MATION	FOR SEQ ID NO: 291:				
	(i)	(A) (B) (C)	NCE CHARACTERISTICS: LENGTH: 336 base pa TYPE: NUCLEIC ACID STRANDEDNESS: DOUBL TOPOLOGY: LINEAR	irs			
	(ii) MOLE	CULE TYPE: CDNA				
	(vi	(A)	INAL SOURCE: ORGANISM: Homo Sapi TISSUE TYPE: Cancer				
	(ix	(B) (C)	URE: NAME/KEY: other LOCATION: 200332 IDENTIFICATION METHO OTHER INFORMATION:		7		
	(ix)	(B) (C)	URE: NAME/KEY: other LOCATION: complemen IDENTIFICATION METHO OTHER INFORMATION:		219		
	(ix	(B) (C)	URE: NAME/KEY: sig_peptic LOCATION: 37330 IDENTIFICATION METHORITHMETHORITHMETHORITHMETHORITHMETHORITHMETHORITHMER INFORMATION:				
	(xi) SEQU	ENCE DESCRIPTION: SE	Q ID NO: 291:			
AAGT	CGCGGT	G GAGC	CAGGCG TGGAAGTCGA CA		AAG GAG ACC Lys Glu Thr -95		54
TAC Tyr	Tyr A	AC ATC sp Ile 90	CTG GGC GTG AAG CCC Leu Gly Val Lys Pro	Ser Ala Ser	CCG GAG AGA Pro Glu Arg	TCA Ser	102

AGA AGG CCT ATC GGA AGC TGG CGC TCA AGT ACC ACC CGG ACA AGA ACC Arg Arg Pro Ile Gly Ser Trp Arg Ser Ser Thr Thr Arg Thr Arg Thr -75 -65

150

CGG Arg -60	ATG Met	AGG Arg	GCG Ala	AGA Arg	AGT Ser -55	TTA Leu	AAC Asn	TCA Ser	TAT Tyr	CCC Pro -50	AGG Arg	CAT His	ATG Met	AAG Lys	TGC Cys -45	198
TTT Phe	CAG Gln	ATC Ile	CAA Gln	AGA Arg -40	AAA Lys	GGG Gly	ATG Met	TTT Phe	ATG Met -35	ACC Thr	AAG Lys	GCG Ala	GAG Glu	AGC Ser -30	AGG Arg	246
CAA G1n	TBV Xaa	AAG Lys	AAG Lys -25	GAG Glu	GCT Ala	CAG Gln	GCA Ala	GCC Ala -20	CCA Pro	GCT Ala	TCT Ser	CTT Leu	CAC His	CCA Pro	TGG Trp	294
ACA Thr	TCT Ser	TTG Leu -10	ACA Thr	TGT Cys	TCT Ser	TTG Leu	GTG Val -5	GTG Val	GTG Val	GAC Asp	GGA Gly	TGC Cys 1	GGG Gly			336

(2) INFORMATION FOR SEQ ID NO: 292:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 397 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 18..194

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 94 region 2..178

id W25476

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 206..359

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100 region 193..346

id W25476

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 359..396

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 94

region 347..384 id W25476

est

(ix) FEATURE:

- (A) NAME/KEY: other(B) LOCATION: 21..278
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97 region 19..276 id HUM179H07B

est

(ix) FEATURE:

- (A) NAME/KEY: other(B) LOCATION: 279..379
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97 region 278..378 id HUM179H07B

est

(ix) FEATURE:

- (A) NAME/KEY: other
 (B) LOCATION: 17..175
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 27..185 id AA002128

est

(ix) FEATURE:

- (A) NAME/KEY: other(B) LOCATION: 171..292
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 182..303 id AA002128

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 358..396
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 373..411 id AA002128

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 325..358
- (C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 339..372 id AA002128

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 204..396
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 186..378

id AA253291

WO 99/06550 PCT/IB98/01232

	EATURE: (A) NAME/KEY: other (B) LOCATION: 22202 (C) FDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 95 region 5185 id AA253291 est	
(EATURE: (A) NAME/KEY: other (B) LOCATION: 42260 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 99 region 26244 id W45609 est	
(EATURE: (A) NAME/KEY: other (B) LOCATION: 251359 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 234342 id W45609 est	
(EATURE: (A) NAME/KEY: other (B) LOCATION: 363396 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 94 region 348381 id W45609 est	
- (EATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 59166 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.8 seq RALSTXLFGSIRG/AA EQUENCE DESCRIPTION: SEQ ID NO: 292:	
AGTGCGCAGA CO	GCAGGGGTC GGCGCCGGGT GAGAGCGTGC GGCCGGATTC ACCACAAC	58
ATG GCA AAT C Met Ala Asn I -35	CTT TTT ATA AGG AAA ATG GTG AAC CCT CTG CTC TAT CTC Leu Phe Ile Arg Lys Met Val Asn Pro Leu Leu Tyr Leu -30 -25	106
AGT CGT CAC A Ser Arg His T -20	ACG GTG AAG CCT CGA GCC CTC TCC ACA NTT CTA TTT GGA Thr Val Lys Pro Arg Ala Leu Ser Thr Xaa Leu Phe Gly -15 -10 -5	154
TCC ATT CGA G	GGT GCA GCC CCC GTG GCT GTG GAA CCC GGG GCA GCA GTG Gly Ala Ala Pro Val Ala Val Glu Pro Gly Ala Ala Val 1 5 10	202

CGC	TCA	CTT	CTC	TCA	CCC	GGC	CTC	CTG	CCC	CAT	CTG	CTG	CCT	GCG	CTG	:	250
															Leu		
		1.5					20					25					

GGG TTC AAA AAC AAG ACT GTC CTT AAG AAG CGC TGC AAG GAC TGT TAC 298 Gly Phe Lys Asn Lys Thr Val Leu Lys Lys Arg Cys Lys Asp Cys Tyr 30 35

CTG GTG AAG AGG CGG GGT CGG TGG TAC GTC TAC TGT AAA ACC CAT CCG 346 Leu Val Lys Arg Arg Gly Arg Trp Tyr Val Tyr Cys Lys Thr His Pro 45 50

AGG CAC AAG CAG AGA CAC ATG TAN ACC CTT TCC CTC CAG AGT CAC GCA 394 Arg His Lys Gln Arg His Met Xaa Thr Leu Ser Leu Gln Ser His Ala 65 70

CAA 397 Gln

(2) INFORMATION FOR SEQ ID NO: 293:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 216 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Cancerous prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 115..216

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97 region 41..142

id H64274

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 74..116

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97 region 1..43

id H64274

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 115..216

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97 region 36..137 id R16956

```
(ix) FEATURE:
```

- (A) NAME/KEY: other
- (B) LOCATION: 79..116
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 1..38

id R16956

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 123..214
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 93

region 57..148

id W04201

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 71..124
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 90

region 4..57

id W04201

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 123..190
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 50..117

id N76590

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 75..116
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 92

region 2..43

id N76590

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(107..195)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 323..411

id N70265

est

(ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: 106..201
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 3.7

seq RIHLCQRSPGSQG/VR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 293:

ACCCTGCCTC ATGCAGCCTA TGGGCTAGGC TTTAGGGTCC GCGGTTGGTC AKACCGGAGC 6

ACTTGGCCTG AAGACCTGGA ATTGGYGACT TCGATATTAA CAAGG ATG GCG GCC 117

Met Ala Ala Ala -30

GCA GCA AGT CGA GGA KTC GGG GCA AAG CTG GGC CTG CGT GAN ATT CGC
Ala Ala Ser Arg Gly Xaa Gly Ala Lys Leu Gly Leu Arg Xaa Ile Arg
-25
-20
-15

ATC CAC TTA TGT CAG CGC TCG CCC GGC AGC CAG GGC GTC AGG GAC TTC 213 Ile His Leu Cys Gln Arg Ser Pro Gly Ser Gln Gly Val Arg Asp Phe -10 -5 1

ATT 216

(2) INFORMATION FOR SEQ ID NO: 294:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 295 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(1..279)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99 region 1..279 id M85423 est
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(196..289)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 466..559 id AA126476

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(133..194)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 560..621 id AA126476 est

(ix) FEATURE:

(A) NAME/KEY: other

(3) LOCATION: complement(105..137)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 90

region 616..648

id AA126476

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 152..292

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 1..141

id R33928

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 160..292

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 14..146

id H67425

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 161..292

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 1..132

id W04820

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 101..232

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 3.7

seq IALTLIPSMLSRA/AG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 294:

AACTTCTTCA TCTTGGTGGT CCTTGCCCAG TTATTTTGCC TCATTAGACA TCAAGAAATG 60

GAGAAAGACT GAAAGTTAAT ATCTTAAGTG CTTGTTCTTC ATG TTT CCT TCT TGT Met Phe Pro Ser Cys

-40

TAT TTA TGC TAT TCT CTT TGT GGC TCC ATT CTT CTT TCA ATC TTC TCA
Tyr Leu Cys Tyr Ser Leu Cys Gly Ser Ile Leu Leu Ser Ile Phe Ser

-35 -30 -25

GCT TAT AAC CGT CTT TCC CTT ATG CTA AGG ATA GCC CTT ACA CTC ATC 211

Ala Tyr Asn Arg Leu Ser Leu Met Leu Arg Ile Ala Leu Thr Leu Ile
-20 -15 -10

CCA TCT ATG CTG TCA AGG GCT GCT GGT TGG TGC TGG TAC AAG GAG CCC

Pro Ser Met Leu Ser Arg Ala Ala Gly Trp Cys Trp Tyr Lys Glu Pro

-5

ACT CAG CAG TTT TCT TAC CTT TGC CTG CCC TGC GGG

Thr Gln Gln Phe Ser Tyr Leu Cys Leu Pro Cys Gly

10 20

(2) INFORMATION FOR SEQ ID NO: 295:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 319 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement (9..318)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98 region 36..345 id R32875 est
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement (52..318)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 35..301 id N69845 est
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement (9..52)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 302..345

id N69845 est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: complement(39..318)
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 46..325 id H20723

(ix)	FEATUR							
		AME/KEY: C						
	(B) L	OCATION: C	complemen	t(3031	8)			
	(0) 0	DENTIFICAT	TON METH					
	(5) 0	THER INFOR	dan 10M:	identit region				
				id HSC3				
				est	011072			
(ix)	FEATUR							
		AME/KEY: c						
	(C) I:	OCATION: c DENTIFICAT	omplemen	t (6531)	8)			
	(D) O	THER INFOR	MATTON .	identity				
	,-, -	211201	22111011.	region 4	_			
				id R021				
				est				
/ : \	CC A MUD	_						
(1X)	FEATUR			-1				
	(B) L(AME/KEY: s OCATION: 1	19_peptio	ae				
	(C) II	DENTIFICAT	ION METHO	DD: Von F	deiine m	.+ ~ i		
	(D) O	THER INFOR	MATION:	score 3.		ICLIX		
				seq QLXE	FLYFVCCIE	PQ/DV		
(CEOUEN	CE DECORED						
(XI)	SEQUENC	CE DESCRIP	TION: SE	2 ID NO:	295:			
						•		
AAMAAGCTCC	CAGCCT	CCAG AGGCT	CTCAA TG	AAGAGTCA	CCTTCATO	GGT CGTC	TYCAGG	60
								00
AACAGGACGG	ATGAMGA	AAGG GGTGG	GGTTA AGA	ACTCAGGG	GCACCTGA	AGG GTCT	GAGCCC	120
CCTT ATG A	GT ACC (CAA GAM GG	A CTG TC	ר אדה ראיז	ר הרם רמנ	ב ככת כתו		1.00
Met S	er Thr (Gln Xaa Gl	y Leu Sei	r Met His	S Ala His	Pro Gl	n Ala	169
-60		- 5	5		-50			
ሞስጥ ክርክ ርር፡	א מייטיים אי	TA MAG GMA	63.6 6.5 -					
TAT ACA CC	n III A. n Phe Ti	lA IAC CIA le Tur Len	His Nia	CGC AAG	AGA CGC	GGA GAG	ATA	217
-45	J 1110 11	-40	nis wid	-35	Arg Arg	GLy Glu		
-							-30	
GGC GAT GC	A GAC TO	CG CGA TTC	AAT GAT	CGA TAT	GCT CAT	AAR AGT	GCT	265
Gly Asp Ala	a Asp Se	er Arg Phe	Asn Asp	Arg Tyr	Ala His	Lys Ser	Ala	203
	-2	25		-20		-15		
CAA TTA TM'	የ ተጥጥ ርባ	ኮር ጥልጥ ጥጥጥ	GTA TCC	ጥርጥ አጥጠ	MMC 033			
CAA TTA TM Gln Leu Xaa	a Phe Le	eu Tvr Phe	Val Cvs	Cve Tla	Pho Cla	GAC GTA	TAT	313
	-10	7	-5	cys iic	rne Gin	Asp var	ryr	
						•		
TAT KTN								319
Tyr Xaa								-
5								
(2) INFORMA	ATION FO	OR SEQ ID 1	NO: 296:					

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 172 base pairs
 (B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens
(F) TISSUE TYPE: Prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(1..170)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99 region 118..287

id AA035134

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(1..170)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99 region 116..285

id N54275

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(1..170)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 119..288 id AA088715

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(19..170)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 115..266

id N78023

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(12..133)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 157..278 id AA100730

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: complement(127..170)
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 90 region 119...1

region 119..162 id AA100730

58

106

154

172

WO 99/06550		322								
		est								
(B) (C)	NAME/KEY: s: LOCATION: 50 IDENTIFICAT:	6118 ION METHOD: V MATION: sco	Von Heijne ma re 3.7 SSCSCSLISFTF							
(xi) SEQUE	ENCE DESCRIPT	TION: SEQ ID	NO: 296:							
ATCTTAGTGC CTTT	ATCTGT CTTTA	TGTCT TGGGGT	TGGG GTAGGTAG	GAT ACCAA ATG Met						
AAA CAC TTT CAG Lys His Phe Gln -20	GAC CTT CCT Asp Leu Pro -15	TCC TCT TGC Ser Ser Cys	AGT TGT TCT Ser Cys Ser -10	TTA ATC TCC Leu Ile Ser -5						
TTT ACT AGA GGA Phe Thr Arg Gly	GAT AAA TAT Asp Lys Tyr 1	TTT GCA TAT Phe Ala Tyr 5	AAT GAA GAA Asn Glu Glu	ATT TTT CTA Ile Phe Leu 10						
GTA TAT AAC GCA Val Tyr Asn Ala 15										
(A) (B)	FOR SEQ ID N CE CHARACTER LENGTH: 424 TYPE: NUCLEI STRANDEDNESS	RISTICS: base pairs C ACID								
(D)	TOPOLOGY: LI	NEAR								

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Hypertrophic prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement(29..369)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 103..443 id W26961

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement(383..424)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97 region 48..89

id W26961

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (34..369)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 100..435

id W26018

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (383..424)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 45..86

id W26018

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (200..369)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 111..280

id W26871

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (143..200)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 281..338

id W26871

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement(383..424)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 56..97 id W26871

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (94..123)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93

region 361..390

id W26871

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (119..369)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 104..354

id W26098

	EATURE: (A) NAME/KEY: other (B) LOCATION: complement(383424) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 4990 id W26098 est
(EATURE: (A) NAME/KEY: other (B) LOCATION: 31302 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 1272 id N99777 est
(CATURE: A) NAME/KEY: other B) LOCATION: 302369 C) IDENTIFICATION METHOD: blastn D) OTHER INFORMATION: identity 95 region 273340 id N99777 est
() ()	ATURE: A) NAME/KEY: sig_peptide B) LOCATION: 155340 C) IDENTIFICATION METHOD: Von Heijne matrix D) OTHER INFORMATION: score 3.7 seq SILGIISVPLSIG/YC
(×i) SE	QUENCE DESCRIPTION: SEQ ID NO: 297:
AGTGAAAAGA AG	ATGCCTAG AGAATGGCAA TTTAAAAGAA AAAGATATAC TTGTTTTGCC 60
	CGACACTG GTTCCCATGA AGCGGCTACC AAAGCTGTTC TCCAGGAGTT 120
TGGTAGAATC GA	CATTCTGG TCAACAATGG TGGA ATG TCC CAG CGT TCT CTG TGC 175 Met Ser Gln Arg Ser Leu Cys -60
ATG GAT ACC AGMet Asp Thr Sc-55	GC TTG GAT GTC TAC AGA RAG CTA ATA GAG CTT AAC TAC 223 er Leu Asp Val Tyr Arg Xaa Leu Ile Glu Leu Asn Tyr -50 -40
TTA GGG ACG G	TG TCC TTG ACA AAA TGT GTT CTG CCT CAC ATG ATC GAG al Ser Leu Thr Lys Cys Val Leu Pro His Met Ile Glu -35 -30 -25
arg bys xaa xa	KA AAG ATT GTT ACT GTG AAT AGC ATC CTG GGT ATC ATA 319 aa Lys Ile Val Thr Val Asn Ser Ile Leu Gly Ile Ile -15 -10
TCT GTA CCT CT Ser Val Pro La	TT TCC ATT GGA TAC TGT GCT AGC RAG CAT GCT CTS HGG 367 eu Ser Ile Gly Tyr Cys Ala Ser Xaa His Ala Leu Xaa

WO 99/06550 PCT/IB98/01232

-5 1 <u>1</u>

GGT TTT TTT AAT RDH CTT CGA ACA GAD CTT GCC ACA TAC CCA GGT ATA
Gly Phe Phe Asn Xaa Leu Arg Thr Xaa Leu Ala Thr Tyr Pro Gly Ile
10 20 25

ATA GTT TCT
Ile Val Ser
424

(2) INFORMATION FOR SEQ ID NO: 298:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 441 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (3) LOCATION: 179..348
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 160..329 id AA159241

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (3) LOCATION: 103..184
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 83..164 id AA159241

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 383..437
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 366..420 id AA159241

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 21..66
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97 region 1..46 id AA159241

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 342..383

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 324..365 id AA159241

est

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 66..102

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 47..83 id AA159241

est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 103..215

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 81..193 id AA076222

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 216..329

(C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 95

region 105 30

region 195..308

id AA076222

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 22..102

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 1..81

id AA076222

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 390..437

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 374..421 id AA076222

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 342..377

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 324..359 id AA076222

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 241..443
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 202..404

id AA149750

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 40..215
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 1..176

id AA149750

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 241..443
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100

region 181..383

id W63593

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 63..184
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 3..124

id W63593

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 179..243
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 92

region 120..184

id W63593

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 320..438
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 267..385

id AA130386

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 216..328
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

328

region 164..276 id AA130386 est

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t	ıх	,	F F.F		UH	۳.	•

(A) NAME/KEY: other

(B) LOCATION: 103..215

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95

region 50..162 id AA130386

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 82..375

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 3.6

seq LALRTSWISSVCS/VT

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 298:

AAG	TGAC	GCG	GCCC	AAGG	GC G	GAAG'	TGAG.	A AA	GTTG	TCTG	CGT	CTCG	AGG	CGAG	TTGGCG	60
GAC	TGTG	CGC	GCGG	CGGG	GC G	ATG Met	GGG Gly	GGC Gly	TCG Ser -95	GGC Gly	AGT Ser	CGC Ar g	CTG Leu	TCC Ser -90	AAG Lys	111
GAG Glu	CTG Leu	CTG Leu	GCC Ala -85	GAR Glu	TAC Tyr	CAG Gln	GAC Asp	TTG Leu -80	ACG Thr	TTC Phe	CTG Leu	ACG Thr	AAG Lys -75	CAG Gln	GAG Glu	159
ATC Ile	CTC Leu	CTA Leu -70	GCC Ala	CAC His	AGG Arg	CGG Arg	TTT Phe -65	TGT Cys	GAG Glu	CTG Leu	CTT Leu	CCC Pro -60	CAG Gln	GAG Glu	CAG Gln	207
CGG Arg	ASK Xaa -55	NGG Xaa	AGT Ser	CGT Arg	CAC His	TTC Phe -50	GGG Gly	CAC His	AAG Lys	TGC Cys	CCT Pro -45	TCG Ser	AGC Ser	AGA Arg	TTC Phe	255
TCA Ser -40	GC€ Ala	TTC Phe	CAG Gln	AGC Ser	TCA Ser -35	AGG Arg	CCA Pro	ACC Thr	CCT Pro	TCA Ser -30	AGG Arg	AGC Ser	GAA Glu	TCT Ser	GCA Ala -25	303
GGG Gly	TCT Ser	TCT Ser	CCA Pro	CAT His -20	CCC Pro	CAG Gln	CCA Pro	AAG Lys	ACA Thr -15	GCC Ala	TTA Leu	GCT Ala	TTG Leu	AGG Arg -10	ACT Thr	351
TCC Ser	TGG Trp	ATC Ile	TCC Ser -5	TCA Ser	GTG Val	TGT Cys	TCA Ser	GTG Val 1	ACA Thr	CAG Gln	CCA Pro	CGC Arg 5	CAG Gln	ACA Thr	TCA Ser	399
AGT Ser	CCC Pro 10	ATT Ile	ATG Met	CCT Pro	TCC Ser	GCA Ala 15	TCT Ser	TTG Leu	ACT Thr	TTG Leu	ATG Met 20	ATG Met	ACG Thr			441

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 284 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 2..162

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99 region 9..169

id N76992

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 162..280

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 168..286

id N76992

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 2..113

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 8..119

id W39234

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 173..280

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 176..283

id W39234

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 113..162

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 118..167

id W39234

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 20..160

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 1..141 id R06371

est

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 193..280

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 173..260

id R06371

est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 159..195

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 94

region 138..174

id R06371

est

(ix) FEATURE:

(A) NAME/KEY: other
(B) LOCATION: 20..159

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 1..140

id R06399

est

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 161..280

(C) IDENTIFICATION METHOD: blastn
(D) OTHER INFORMATION: identity 97

region 141 20

region 141..260

id R06399

est

(ix) FEATURE:

(A) NAME/KEY: other(B) LOCATION: 27..165

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 1..139

id AA043154

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 166..280

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 99

region 141..255

id AA043154

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

60

WO 99/003	30	. 331										
(B) LOCATION: 132215(C) IDENTIFICATION METHOD: Von Heijne matrix(D) OTHER INFORMATION: score 3.6seq PLSDSWALLPASA/GV												
(xi)	SEQUI	ENCE DE	ESCRIPTION:	SEQ ID NO:	299:							
AACAACTTCC	GGCC	CCACTG	AGCGGTGTCC	TGAGCCGATT	ACAGCTAGGT	AGTGGAGCGC						
CGCTGCTTAC	CTGG	GTGCAG	GAGACAGCCG	GAGTCGCTGG	GGGAGCTCCG	CGCCGCCGGA						

CGCTGCTTAC TCCG CGCCGCCGGA 120 CGCCCGTGAC C ATG TGG AGG CTG CTG GCT CGC GCT AGT GCG CCG CTC CTG 170 Met Trp Arg Leu Leu Ala Arg Ala Ser Ala Pro Leu Leu -25 CGG GTG CCC TTG TCA GAT TCC TGG GCA CTC CTC CCC GCC AGT GCT GGC 218 Arg Val Pro Leu Ser Asp Ser Trp Ala Leu Leu Pro Ala Ser Ala Gly -15 -10 -5 GTA AAG ACA CTG CTC CCA GTA CCA AGT TTT GAA GAT GTT TCC ATT CCT 266 Val Lys Thr Leu Leu Pro Val Pro Ser Phe Glu Asp Val Ser Ile Pro 10 15 GAA AAA CCC AAG CTA CTG 284 Glu Lys Pro Lys Leu Leu 20

(2) INFORMATION FOR SEQ ID NO: 300:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 374 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 169..332
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 163..326 id H71676

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 92..170
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 94 region 87..165

id H71676

	(ix)	(A) (B) (C)	NAM LOC I DE	E/KE ATIO NTIF	N: 2	ther 08 ION MATI	5 METH	ide reg	ntit ion H716	y 93 18					
	(ix)	(A) (B) (C)	NAM LOC.	ATIO NTIF	N: 3 ICAT	ther 34 ION MATI	METH	ide reg	blas ntit ion H716	y 93 330.					
	(.	ix)	(A) (B) (C)	NAM! LOCA	ATIO	N: 2	ther 64 ION I	METH	ide: reg:	olas ntit ion :	y 96 31	15				
		ix) ki)	(A) (B) (C) (D)	NAME LOCA I DEN	ATION NTIF: ER IN	N: 6 [CAT] NFORM	ig_pe 34 [*] ION N 4ATIO	7 METHO DN:	DD: V scor seq	re 3. ATFV	. 6 /TQA]	LIQX				
AAA	AA A1 Me	TG GG	CG GA	AT CA	is Va	TG CA al GI	AG AG ln Se	GC C	TG GG	la G	AA C' ln Lo 105	TA GA	AG A	AT C	TG TGC eu Cys -100	50
AAA Lys	CAG Gln	Leu	TAT Tyr	Glu	Thr	ACA Thr	GAC Asp	ACA Thr	RSC Xaa -90	AST Xaa	CGG Arg	AGC Ser	TCC Ser	SAG Xaa -85	GCA Ala	98
GAG Glu	AAA Lys	GCS Ala	TTG Leu -80	GTT Val	GAR Glu	TTT Phe	ACC Thr	AAC Asn -75	AGC Ser	CCT Pro	GAT Asp	TGC Cys	CTG Leu -70	AGC Ser	AAG Lys	146
TGC Cys	CAG Gln	CTA Leu -65	CTC Leu	CTC Leu	GAA Glu	AGA Arg	GGA Gly -60	AGT Ser	TCC Ser	TCT Ser	TAC Tyr	TCC Ser -55	CAG Gln	TTA Leu	CTG Leu	194
GCA Ala	GCT Ala -50	ACA Thr	TGC Cys	CTT Leu	ACC Thr	AAG Lys -45	CTT Leu	GTA Val	TCA Ser	Arg	ACA Thr -40	AAC Asn	AAC Asn	CCC Pro	CTA Leu	242
CCA Pro	TTG Leu	GAA Glu	CAG Gln	CGA Arg	ATA Ile	GAT Asp	ATT Ile	CGG Arg	AAC Asn	TAT Tyr	GTG Val	CTC Leu	AAC Asn	TAS Xaa	CTT Leu	290

GCC ACT CGG CCG AAG TTG GCT ACT TTC GTG ACA CAA GCA CTT ATT CAG Ala Thr Arg Pro Lys Leu Ala Thr Phe Val Thr Gln Ala Leu Ile Gln -15

TKA TAT GCC AGA ATC ACA AAA CTG GGC TGG TTT GAC Arg Tyr Ala Arg Ile Thr Lys Leu Gly Trp Phe Asp

(2) INFORMATION FOR SEQ ID NO: 301:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 238 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 15..235
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99 region 2..222 id H39781

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 16..173
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 100 region 1..158 id AA017398

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 172..235
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 98

region 159..222

id AA017398

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 16..235
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 1..220

id AA059110

	(A) NAME/KEY: other(B) LOCATION: 17235(C) IDENTIFICATION METH(D) OTHER INFORMATION:	OD: blastn identity 99 region 1219 id AAO37143 est	
	EATURE: (A) NAME/KEY: other (B) LOCATION: 56235 (C) IDENTIFICATION METHO (D) OTHER INFORMATION:	DD: blastn identity 99 region 56235 id R75754 est	
(A) NAME/KEY: other B) LOCATION: 1455 C) IDENTIFICATION METHO D) OTHER INFORMATION:	DD: blastn identity 100 region 1556 id R75754 est	
((A) NAME/KEY: sig_peptid B) LOCATION: 62226 C) IDENTIFICATION METHO D) OTHER INFORMATION:	D: Von Heijne matrix score 3.6 seq TCSVCCYLFWLIA/IP	
		GGTAGGG GTTGGCGCTC AGGCGGCGAC	
C ATG GCG TAT	CAC GGC CTC ACT GTG CC	T CTC ATT GTG ATG AGC GTG TTC to Leu Ile Val Met Ser Val Phe -45 -40	109
TGG GGC TTC G Trp Gly Phe V	al Gly Phe Leu Val Pro	TGG TTC ATC CCT AAG GGT CCT Trp Phe Ile Pro Lys Gly Pro -30 -25	157
Asn Arg Gly V	TT ATC ATT ACC ATG TTG al Ile Ile Thr Met Leu -15	GTG ACC TGT TCA GTT TGC TGC Val Thr Cys Ser Val Cys Cys -10	205
TAT CTC TTT T Tyr Leu Phe T -5	GG CTG ATT GCA ATT CCG rp Leu Ile Ala Ile Pro	GCC TGG Ala Trp	238
(2) INFORMATI	ON FOR SEQ ID NO: 302:		

(ix) FEATURE:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 437 base pairs

335 . (B) TYPE: NUCLEIC ACID (C) STRANDEDNESS: DOUBLE (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: CDNA (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Cancerous prostate (ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: complement (397..432) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 97 region 283..318 id H83411 est (ix) FEATURE: (A) NAME/KEY: sig peptide (B) LOCATION: 54..227 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.5 seq GGILMGSFQGTIA/GO (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 302: ATATTTGCCC CTTACTTTAT CTTGTGCCTT GAGAAATTGC TGGGGAGAGA GGT ATG 56 Met TCC ACT GGG CAG CTG TAC AGG ATG GAG GAT ATA GGG CGT TTC CAC TCC Ser Thr Gly Gln Leu Tyr Arg Met Glu Asp Ile Gly Arg Phe His Ser -55 -50 CAG CAG CCA GGT TCC CTC ACC CCA AGC TCA CCC ACT GTT GGG GAG ATT 152 Gla Gla Pro Gly Ser Leu Thr Pro Ser Ser Pro Thr Val Gly Glu Ile -35 ATC TAC AAT AAC ACC AGA AAC ACA TTG GGG TGG ATT GGG GGT ATC CTT 200 Ile Tyr Asn Asn Thr Arg Asn Thr Leu Gly Trp Ile Gly Gly Ile Leu -20 -15 ATG GGT TCT TTT CAG GGA ACC ATT GCT GGA CAA GGC ACA GGA GCC ACC Met Gly Ser Phe Gln Gly Thr Ile Ala Gly Gln Gly Thr Gly Ala Thr TCC ATT TCT GAG CTC TGC AAG GGA CAA GAA CTA GAG CCA TCA GGG GCT 296 Ser Ile Ser Glu Leu Cys Lys Gly Gln Glu Leu Glu Pro Ser Gly Ala 10 GGG CTC ACT GTG GCC CCA CCC CAA GCC GTC AGC CTC CAG GGA TCA CAC Gly Leu Thr Val Ala Pro Pro Gln Ala Val Ser Leu Gln Gly Ser His

CCT GCC TTG GCT GCT ACA GCT TTT TCA CTC CAS TGC CCT AGG GGA GTT

Pro Ala Leu Ala Ala Thr Ala Phe Ser Leu Xaa Cys Pro Arg Gly Val

CAG CAS CTA ATG ATC TCT ATC TCT GAA CAT CTC TTC ATC CAT GCT

392

437

-40

-25

Gln Xaa Leu Met Ile Ser Ile Ser Glu His Leu Phe Ile His Ala 60 65 70

(2) INFORMATION FOR SEQ ID NO: 303:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 353 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 27..347
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 1..321 id T31485

-u 151

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 42..352
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99

region 1..311

id HSC38B061

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 135..325
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 97

region 70..260

id T66273

est

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 69..140
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 93

region 5..76

id T66273

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 2..220
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 96 region 6..224

337

id R24829 est

(ix)	FEATURE:
------	----------

(A) NAME/KEY: other(B) LOCATION: 236..275

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 100

region 243..282

id R24829

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 50..318

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 98

region 10..278 id HSC2LF071

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 282..332

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 3.5

seq RWWCFHLQAEASA/HP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 303:

ATAATAT CTAAAAAGCT AAATTTTAAA TACCAGCTTT ACATAAATGA TTGTKGACTC 60

TGGTCTGTKT CTGACACCTT TCCAGAAAAA AGTCAATTGT TCAGGTACAC CAAAGAGGAA 120

GAAGAGCTGT GGAGGCCACC CTCTACAAAG CTTTATAGAA CTTCTGGATC TAACTCACAA 180

ACAAGCTTCC AGAAGAGACT AGAGACCTTA GGCCAGGAGA TGAAGGAGTT CAGTAGCAAA 240

GTCACACCTG TCCAATTCCC TGAGCTTTGC TCACTCAGCT A ATG GGA TGG CAA AGG 296

Met Gly Trp Gln Arg

-15

TGG TGG TGC TTT CAT CTT CAG GCA GAA GCC TCT GCC CAT CCC CCT CAA
Trp Trp Cys Phe His Leu Gln Ala Glu Ala Ser Ala His Pro Pro Gln

-10 -5 1

GGG CTG CAG

353

Gly Leu Gln

(2) INFORMATION FOR SEQ ID NO: 304:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 260 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) (ORIGINAL SOURCE: (A) ORGANISM: Homo Sapie (F) TISSUE TYPE: Hypertr	ens cophic prostate	
(xi)	FEATURE: (A) NAME/KEY: other (B) LOCATION: 80236 (C) IDENTIFICATION METHO (D) OTHER INFORMATION:	DD: blastn identity 95 region 34190 id N34164 est	
(ix) E		D: blastn identity 98 region 66232 id R89543 est	
		D: blastn identity 98 region 66229 id H59647 est	
	EATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 126170 (C) IDENTIFICATION METHOR (D) OTHER INFORMATION:	D: Von Heijne matrix	
(xi) S	EQUENCE DESCRIPTION: SEQ	ID NO: 304:	
		AGATTGC TGAGGAGGCG GCGGGTAGCT	60
GGCAGGCGCC G	ACTTCCGAA GGCCGCCGTC CGG	GCGAGGT GTCCTCATGA CTTCTCTTGT	120
GGACC ATG TC Met Se -15	C GTG ATC TTT TTT GCC TG r Val Ile Phe Phe Ala Cy -10	C GTG GTA CGG GTA AGG GAT GGA s Val Val Arg Val Arg Asp Gly -5	170
CTG CCC CTC Leu Pro Leu	TCA GCC TCT ACT GAT TTT Ser Ala Ser Thr Asp Phe 5	TAC CAC ACC CAA GAT TTT TTG Tyr His Thr Gln Asp Phe Leu 10	218
GAA TGG AGG . Glu Trp Arg .	AGA CGG CTC AAG AGT TTA (Arg Arg Leu Lys Ser Leu 1 20 25	GCC TTG CGA CTG AAG Ala Leu Arg Leu Lys 30	260

(2) INFORMATION FOR SEQ ID NO: 305:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 242 base pairs

(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 44..210

. (C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 92

region 29..195

id R88607

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 17..135

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 16..134 id AA035300

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 136..244

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 134..242

id AA035300

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 38..244

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 1..207

id AA147873

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (128..244)

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 96

region 263..379

id AA147836

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: complement (38131) (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 94 region 375468 id AA147836 est	
<pre>(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 136244 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 95</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 45138 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 95</pre>	
<pre>(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 66113 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.5</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 305:	
AATTAGCGCG TAACGCASAG ACTGCTTGCT GCGGCAGAGA CGCCAGAKGT GCAGCTCCAG	60
CAGCA ATG GCA GTG ACG GCG TTG GCG GCG MRG ACG TGG CTT GGC GTG TGG Met Ala Val Thr Ala Leu Ala Ala Xaa Thr Trp Leu Gly Val Trp -15 -10 -5	110
GGC GTG AGG ACC ATG CAA GCC CGA GGC TTC GGC TCG GAT CAG TCC GAG Gly Val Arg Thr Met Gln Ala Arg Gly Phe Gly Ser Asp Gln Ser Glu 1 5 10 15	158
AAT GTC GAC CGG GGC GCG GGC TCC ATC CGG GAA GCC GGT GGG GCC TTC Asn Val Asp Arg Gly Ala Gly Ser Ile Arg Glu Ala Gly Gly Ala Phe 20 25 30	206
GGA AAS AGA GAG CAG GCT GAA GAS SAA CGA TAT TTC Gly Xaa Arg Glu Gln Ala Glu Xaa Xaa Arg Tyr Phe 35 40	242
(2) INFORMATION FOR SEQ ID NO: 306:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 402 base pairs(B) TYPE: NUCLEIC ACID(C) STRANDEDNESS: DOUBLE	

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 151..402
- (C) IDENTIFICATION METHOD: fasta

(D) OTHER INFORMATION: identity 100.0 region 1..252 id HSU21128

vrt

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 155..402
- (C) IDENTIFICATION METHOD: fasta
- (D) OTHER INFORMATION: identity 99.6 region 1..248 id HSU18728 vrt

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 131..402
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100 region 1..272 id H27256 est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 161..402
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 31..272 id W95921

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 296..402
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 100 region 141..247

id C17793

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 151..252
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 1..102

id C17793

	EATURE: (A) NAME/KEY: other (B) LOCATION: 174402 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 1229 id AA180902 est
	EATURE: (A) NAME/KEY: other (B) LOCATION: 199402 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 1204 id R58323 est
·	CATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 235288 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 12 seq FTLFLALIGGTSG/QY
(xi) S	QUENCE DESCRIPTION: SEQ ID NO: 306:
ACATGCCACA CO	CACAAGATC CCCACAATGA CATAACTCCA TTCAGAGACT GGCGTGACTG 60
	CCACCCC CCCTTCAGCT CTTGTATCAC TCAGAATCTG GCAGCCAGTT 120
	SAGTTCACA GCATATATTG GTGGATTCTT GTCCATAGTG CATCTGCTTT 180
	AAGCAGTG TCAAGACAGT AAGGATTCAA ACCATTTGCC AAAA ATG 237 Met
-15	CA TTT ACT CTC TTC CTG GCA TTG ATT GGT GGT ACC AGT 285 Lia Phe Thr Leu Phe Leu Ala Leu Ile Gly Gly Thr Ser -10 -5
Gly Gin Tyr 1	AT GAT TAT GAT TTT CCC CTA TCA ATT TAT GGG CAA TCA 'yr Asp Tyr Asp Phe Pro Leu Ser Ile Tyr Gly Gln Ser 10 15
TCA CCA AAC I Ser Pro Asn C	GT GCA CCA GAA TGT AAC TGC CCT GAA AGC TAC CCA AGT ys Ala Pro Glu Cys Asn Cys Pro Glu Ser Tyr Pro Ser 20 25 30
	GT GAT GAG CTG ys Asp Glu Leu 35

(2) INFORMATION FOR SEQ ID NO: 307:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 274 base pairs

(B) TYPE: NUCLEIC ACID

WO 99/06550 PCT/IB98/01232 343

(C) STRANDEDNESS: DOUBLE

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 120..272

(C) IDENTIFICATION METHOD: fasta

(D) OTHER INFORMATION: identity 96.1 region 1..151 id HSU21128 vrt

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 124..272

(C) IDENTIFICATION METHOD: fasta

(D) OTHER INFORMATION: identity 96.0 region 1..147 id HSU18728

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 141..272

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97 region 40..171 id H27256

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 100..136

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97

region 1..37 id H27256

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 141..272

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 97 region 40..171

id W95921

est

(ix) FEATURE:

(A) NAME/KEY: other

(B) LOCATION: 141..245

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 95 region 52..156 id AA093526

344	
est	
(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 89136 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 95 region 249 id AA093526 est	
(ix) FEATURE:	
(A) NAME/KEY: other (B) LOCATION: 145272 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98 region 1128 id AA180902 est	
(ix) FEATURE:	
(A) NAME/KEY: other (B) LOCATION: 141223 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 93 region 20102 id C17793 est	
(ix) FEATURE:	
(A) NAME/KEY: sig_peptide (B) LOCATION: 206259 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 12	
seq FTLFLALIGGTSG/QY	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 307:	
ATAACTCCAT TCAGAGACTG GCGTGACTGG GCTGGGTCTC CCCACCCCCC CCTTCAGCTC	60
TTGTATGACT CAGAATCTGG CAGCCAGTTC CGTCCTGACA GAGTTCACAG CATATATTGG	120
TGGATTCTTG TCCAWAAGTG GVATCTGCTT TARGAWTTAA CGAAAGCAGT GTCAAGACAG	180
TAAGGATTCA AACCATTTGC CAAAA ATG AGT CTA AGT GCA TTT ACT CTC TTC Met Ser Leu Ser Ala Phe Thr Leu Phe -15 -10	232
CTG GCA TTG ATT GGT GGT ACC AGT GGC CAG TAC TAT GAT TGG Leu Ala Leu Ile Gly Gly Thr Ser Gly Gln Tyr Tyr Asp Trp -5 1 5	274

(2) INFORMATION FOR SEQ ID NO: 308:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 436 base pairs
(B) TYPE: NUCLEIC ACID

(C) STRANDEDNESS: DOUBLE

PCT/IB98/01232 WO 99/06550 345

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Hypertrophic prostate

(ix) FEATURE:

- (A) NAME/KEY: other (B) LOCATION: 65..433
- (C) IDENTIFICATION METHOD: fasta
- (D) OTHER INFORMATION: identity 100.0 region 1..369 id HSU21128

vrt

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 69..433
- (C) IDENTIFICATION METHOD: fasta
- (D) OTHER INFORMATION: identity 99.7 region 1..365 id HSU18728 vrt

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 45..433
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99 region 1..389 id H27256

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 75..433
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 31..389 id W95921

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 210..433
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 141..364

id C17793

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 65..166
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98 region 1..102 id C17793

(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 88433 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 100 region 1346 id AA180902 est												
<pre>(ix) FEATURE: (A) NAME/KEY: other (B) LOCATION: 113370 (C) IDENTIFICATION METHOD: blastn (D) OTHER INFORMATION: identity 98</pre>												
(ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: 149202												
(C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 12 seq FTLFLALIGGTSG/QY												
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 308:												
AGCTCTTGTA TCACTCAGAA TCTGGCAGCC AGTTCCGTCC TGACAGAGTT CACAGCATAT 6												
ATTGGTGGAT TCTTGTCCAT AGTGCATCTG CTTTAAGAAT TAACGAAAGC AGTGTCAAGA	A 120											
CAGTAAGGAT TCAAACCATT TGCCAAAA ATG AGT CTA AGT GCA TTT ACT CTC Met Ser Leu Ser Ala Phe Thr Leu -15	172											
TTC CTG GCA TTG ATT GGT GGT ACC AGT GGC CAG TAC TAT GAT TAT GAT Phe Leu Ala Leu Ile Gly Gly Thr Ser Gly Gln Tyr Tyr Asp Tyr Asp -10 -5 1 5	220											
TTT CCE CTA TCA ATT TAT GGG CAA TCA TCA CCA AAC TGT GCA CCA GAA Phe Pro Leu Ser Ile Tyr Gly Gln Ser Ser Pro Asn Cys Ala Pro Glu 10 15 20	268											
TGT AAC TGC CCT GAA AGC TAC CCA AGT GCC ATG TAC TGT GAT GAG CTG Cys Asn Cys Pro Glu Ser Tyr Pro Ser Ala Met Tyr Cys Asp Glu Leu 25 30 35	316											
Cys Asn Cys Pro Glu Ser Tyr Pro Ser Ala Met Tyr Cys Asp Glu Leu	316											
Cys Asn Cys Pro Glu Ser Tyr Pro Ser Ala Met Tyr Cys Asp Glu Leu 25 30 35 AAA TTG AAA AGT GTA CCA ATG GTG CCT CCT GGA ATC AAG TAT CTT TAC Lys Leu Lys Ser Val Pro Met Val Pro Pro Gly Ile Lys Tyr Leu Tyr												

(2)	INFO)RMA1	LION	FOR	SEQ	ID t	10: 3	309:								
	(i	.) SE	(B) (C)	LENG TYPE STRA	CHARA TH: C: NU NDED DLOGY	423 CLEI NESS	base C AC	e pai CID OUBLE								
	(i	i) M	OLEC	ULE	TYPE	: CE	NA									
	(v	·i) C		ORGA	SOUR NISM UE T	1: Ho				tate	:					
	(i	x) E	(B)	NAME LOCA IDEN	KEY TION TIFI R IN	: 75 CATI	34 ON M	IETHO N:	iden regi	tity on 1	96. 26 BAO7M	9				
	(i	x) E	(B) (C)	NAME LOCA I DEN	KEY TION TIFI R IN	: 51 CATI	15 ON M	IETHC)N:	iden	tity on 1	, 99 10	9				
	(i	×) F	(B) (C)	NAME LOCA IDEN	/KEY TION TIFI R IN	: 91 CATI	15 ON M	O ETHO	D: V	e 8.	leijn 9 LLPF					
	(x	i) S	EQUE	NCE	DESC	RIPT	'ION:	SEÇ) ID	NO:	309:					
AATT	TTGAA	TT C	GGGG	GTGT	C TA	\GAA#	\GAGA	A AGO	CATA	AGTC	GGCG	AGC#	AC C	CTG	AGCAT	60
CCC	GCTCI	GG 1	rgcco	CTG	CA GC	CGGC	CAGAC		: Val					Pro	CTG Leu	114
	CTC Leu															162
AGG Arg 5	AAA Lys	ATG Met	CTG Leu	TCC Ser	AGT Ser 10	GGG Gly	GTG Val	TGT Cys	ACA Thr	TCA Ser 15	ACT Thr	GTT Val	CAG Gln	CTT Leu	CCT Pro 20	210
GGG Gly	AAA Lys	GTA Val	GTT Val	GTG Val 25	GTC Val	ACA Thr	GGA Gly	GCT Ala	AAT Asn 30	ACA Thr	GGT Gly	ATC Ile	GGG Gly	AAG Lys 35	GAG Glu	258

ACA Thr	GCC Ala	AAA Lys	GAG Glu 40	CTG Leu	GCT Ala	CAG Gln	AGA Arg	GGA Gly 45	GCT Ala	CGA Arg	GTA Val	TAT Tyr	KTA Xaa 50	GCT Ala	TNN Xaa	306
NGG Xaa	GAT Asp	GTG Val 55	GAA Glu	AAG Lys	GGG Gly	GAA Glu	TTG Leu 60	GTG Val	GCC Ala	ARA Xaa	GAG Glu	ATC Ile 65	CAG Gln	ACC Thr	ACG Thr	354
ACA Thr	GGG Gly 70	AAN Xaa	SAG Xaa	CAG Gln	GTG Val	TTG Leu 75	GTG Val	CGG Arg	RAA Xaa	CTG Leu	GAC Asp 80	CTG Leu	TCT Ser	GAT Asp	ACT Thr	402
	TCT Ser						•									423

(2) INFORMATION FOR SEQ ID NO: 310:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 306 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 132..303
 - (C) IDENTIFICATION METHOD: fasta
 - (D) OTHER INFORMATION: identity 96 region 1..171

id HSClR

vrt

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 143..303
 - (C) IDENTIFICATION METHOD: fasta
 - (D) OTHER INFORMATION: identity 98

region 24..183

id HUMC1R

vrt

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 181..303
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 92

region 1..123

id T74375

est

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 170..213

(C) IDENTIFICATION METHOD: blastn

(D) OTHER INFORMATION: identity 93

region 1..44 id T64778

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 184..228

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 8.1

seq LLYLLVPALFCRA/GG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 310:

AAAAACTCAG ATCTTTTGTT TATGCAAATA GTTCATTCCC TCCAACATTC CTCCGGGAAT 60

GGTCCCCCCT CCACTCCACA GAAAACCCTC CCCTCCCTGC TGTGCATGAC GCGGGCTCCC 120

TCTGSACACA GKGVMCRAAG ACGCTGTCGG GAKAGCCCCA GGATTCAACA CGGGCCTTGA 180

GAA ATG TGG CTC TTG TAC CTC CTG GTG CCG GCC CTG TTC TGC AGG GCA 228

Met Trp Leu Leu Tyr Leu Leu Val Pro Ala Leu Phe Cys Arg Ala

-15

-10

-5

GGA GGC TCC ATT CCC ATC CCT CAG AAG TTA TTT GGG GAG GTG ACT TCC

Gly Gly Ser Ile Pro Ile Pro Gln Lys Leu Phe Gly Glu Val Thr Ser

CCT CTG TTC CCC AAG CCT TAC CCC AAC GGG
Pro Leu Phe Pro Lys Pro Tyr Pro Asn Gly
20 25

(2) INFORMATION FOR SEQ ID NO: 311:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 263 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 50..263
 - (C) IDENTIFICATION METHOD: fasta
 - (D) OTHER INFORMATION: identity 99 region 1..214 id HSSPG28 vrt

	(ix)	(B) (C)	NAM LOC I DE	E/KE ATIO NTIF ER I	N: 7 ICAT	52 ION	63 METH	ide reg	fast ntit ion HSCR	y 99 11	89				
	(ix)	(B) (C)	NAMI LOCA I DEI	E/KEY ATION NTIF: ER IN	N: 5: ICAT:	$1.\overline{.}1$	46 METHO	DD: 1	re 7	. 7	ne ma				
	(:	xi)	SEQUI	ENCE	DESC	CRIP	rion	: SE	DI C	NO:	311	:				
AAT	АТАТ	ACG	GCTC'	TAAC	CT TO	CTCT	CTCT(G CA	CCT T (CCTT	CTG	TCAA'		ATG A		56
CAA Gln -30	ATA Ile	CTT Leu	CAT His	CCT Pro	GCT Ala -25	CTG Leu	GAA Glu	ACC Thr	ACT Thr	GCA Ala -20	ATG Met	ACA Thr	TTA Leu	TTC Phe	CCA Pro -15	104
GTG Val	CTG Leu	TTG Leu	TTC Phe	CTG Leu -10	GTT Val	GCT Ala	GGG Gly	CTG Leu	CTT Leu -5	CCA Pro	TCT Ser	TTT Phe	CCA Pro	GCA Ala 1	AAT Asn	152
GAA Glu	GAT Asp	AAG Lys 5	GAT Asp	CCC Pro	GCT Ala	TTT Phe	ACT Thr 10	GCT Ala	TTG Leu	TTA Leu	ACC Thr	ACC Thr 15	CAA Gln	ACA Thr	CAA Gln	200
GTG Val	CAA Gln 20	AGG Arg	GAG Glu	ATT Ile	GTG Val	AAT Asn 25	AAG Lys	CAC His	AAT Asn	GAA Glu	CTG Leu 30	AGG Arg	AGA Arg	GCA Ala	GTA Val	248
			GCC Ala													263
(2)			rion EQUEN													
	, 1	., 5.	(A) (B) (C)	LENG TYPE STRA	TH: : NU NDED LOGY	465 CLEI NESS	base C AC : DO	pai ID UBLE								
	(i	i) N	OLEC	ULE	TYPE	: CD	NA									
	(v	i) (RIGI (A) (F)	ORGA	SOUR NISM UE T	: Ho	mo S Can	apie cero	ns us p	rost	ate					
	(i	×) Ē	FEATU (A) (B)	NAME	/KEY TION			67								

id HSU03877

vrt

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 213..467
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 232..486

id AA150097

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 35..204
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 55..224

id AA150097

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 43..467
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 56..480

id AA155808

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 43..404
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 73..434

id AA147966

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 395..467
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 424..496

id AA147966

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 51..467
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 98

region 1..417

id AA058479

est

(ix) FEATURE:

		JJ2
	NAME/KEY: other	
(B)	LOCATION: 70405	
	IDENTIFICATION MET	
(D)	OTHER INFORMATION	: identity 97 region 1336 id W46890 est
EAT	URE:	
(A)	NAME/KEY: other	
(B)	LOCATION: 394425	5
(C)	IDENTIFICATION MET	THOD: blastn

(ix) FEATURE:

(C) IDENTIFICATION METHOD: blastn
(D) OTHER INFORMATION: identity 100 region 326..357 id W46890

est

(ix) FEATURE:

(A) NAME/KEY: sig_peptide (B) LOCATION: 52..102

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 6.9

seq LFLTMLTLALVKS/QD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 312:

AACTC	CCCTC	GCTG	CCCG	GG C	CCGG	AGCG	C AS:	SNGG	CCGC	ACA	GATT:	CAC		G TTG t Leu	57
AAA GG Lys A:	CC CTT la Leu	TTC Phe	CTA Leu	ACT Thr -10	ATG Met	CTG Leu	ACT Thr	CTG Leu	GCG Ala -5	CTG Leu	GTC Val	AAG Lys	TCA Ser	CAG Gln 1	105
GAC AC	CC GAA nr Glu	GAA Glu 5	ACC Thr	ATC Ile	ACG Thr	TAC Tyr	ACG Thr 10	CAA Gln	T GC Cys	ACT Thr	GAC Asp	GGA Gly 15	TAT Tyr	GAG Glu	153
TGG GA	AT CCT sp Pro 20	GTG Val	AGA Arg	CAG Gln	CAA Gln	TGC Cys 25	AAA Lys	GAT Asp	ATT Ile	GAT Asp	GAA Glu 30	TGT Cys	GAC Asp	ATT Ile	201
GTC CC Val Pr	CA GAC co Asp 35	GCT Ala	TGT Cys	AAA Lys	GGT Gly 40	GGA Gly	ATG Met	AAG Lys	TGT Cys	GTC Val 45	AAC Asn	CAC His	TAT Tyr	GGA Gly	249
GGA TA Gly Ty 50	AC CTC /r Leu	TGC Cys	CTT Leu	CCG Pro 55	AAA Lys	ACA Thr	GCC Ala	CAG Gln	ATT Ile 60	ATT Ile	GTC Val	AAT Asn	AAT Asn	GAA Glu 65	297
CAG CC Gln Pr	CT CAG	CAG Gln	GAA Glu 70	ACA Thr	CAA Gln	CCA Pro	GCA Ala	GAA Glu 75	GGA Gly	ACC Thr	TCA Ser	GGG Gly	GCA Ala 80	ACC Thr	345
ACC GC Thr Gl	GG GTT Ly Val	GTA Val 85	GCT Ala	GCC Ala	DNC Xaa	AGC Ser	ATG Met 90	GCA Ala	ACC Thr	AGT Ser	GBA Xaa	GTG Val 95	TTG Leu	MNN Xaa	393
GGG GG	GT GGT y Gly 100	TTT Phe	GTG Val	GCC Ala	AGT Ser	GCT Ala 105	GCT Ala	GCA Ala	GTC Val	GCA Ala	GGC Gly 110	CCT Pro	GAA Glu	ATG Met	441

WO 99/06550 PCT/IB98/01232

CAG ACT GGC CGG AAT AAC TTT GTC Gln Thr Gly Arg Asn Asn Phe Val 115

465

(2) INFORMATION FOR SEQ ID NO: 313:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 256 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 50..256
 - (C) IDENTIFICATION METHOD: fasta
 - (D) OTHER INFORMATION: identity 96 region 1..204 id HUMTCAYV

vrt

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 50..256
 - (C) IDENTIFICATION METHOD: fasta
 - (D) OTHER INFORMATION: identity 93 region 1..207 id MACTCRAAQ

vrt

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 50..256
 - (C) IDENTIFICATION METHOD: fasta
 - (D) OTHER INFORMATION: identity 94

region 1..204 id MACTCRAAR

vrt

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: 50..115
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq LLILWFHLDCVSS/IL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 313:

AAT Asn	CCT Pro	TTG Leu	GCA Ala	GCC Ala -15	CCA Pro	TTA Leu	CTA Leu	ATC Ile	CTC Leu -10	TGG Trp	TTT Phe	CAT His	CTT Leu	GAC Asp -5	TGC Cys	106
GTG Val	AGC Ser	AGC Ser	ATA Ile 1	CTG Leu	AAC Asn	GTG Val	GAA Glu 5	CAA Gln	AGT Ser	CCT Pro	CAG Gln	TCA Ser 10	CTG Leu	CAT His	GTT Val	154
CAG Gln	GAG Glu 15	GGA G1 y	GAC Asp	AGC Ser	ACC Thr	AAT Asn 20	TTC Phe	ACC Thr	TGC Cys	AGC Ser	TTC Phe 25	CCT Pro	TCC Ser	AGC Ser	AAT Asn	202
TTT Phe 30	TAT Tyr	GCC Ala	TTA Leu	CAC His	TGG Trp 35	TAC Tyr	AGA Arg	TGG Trp	GAA Glu	ACT Thr 40	GCA Ala	AAA Lys	AGC Ser	CCC Pro	GAG Glu 45	250
GCC Ala																256

(2) INFORMATION FOR SEQ ID NO: 314:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 455 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 62..455
 - (C) IDENTIFICATION METHOD: fasta
 - (D) OTHER INFORMATION: identity 98.7 region 1..392 id HSU32907

vrt

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 138..415
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 99 region 1..278 id H09504

- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (B) LOCATION: 410..454
 - (C) IDENTIFICATION METHOD: blastn
 - (D) OTHER INFORMATION: identity 91 region 274..318 id H09504

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 160..455
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 1..296 id H17686

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 128..329
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 42..243 id AA247900

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 85..123
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 94

region 1..39 id AA247900

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 318..355
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 231..268 id AA247900

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 128..231
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 96

region 22..125

id R57541

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 231..274
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 97

region 124..167

id R57541

est

(ix) FEATURE:

- (A) NAME/KEY: other
- (B) LOCATION: 312..455
- (C) IDENTIFICATION METHOD: blastn
- (D) OTHER INFORMATION: identity 99

region 1..144 id N87278 est

,	iх	١	FEATURE:	
l	1 X	1	LEATURE:	۰

(A) NAME/KEY: sig_peptide

(B) LOCATION: 345..389

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.3

seq VVTIVILLCFCKA/AE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 314:

AGCTGGGGC	ATGTAATTTA 2	AAACCTCTCA	ስ ስ ስ ርጥር ጥር ር ጥ	GCGGTCCGTG CACAGCATTA	
					60
GTATAACGT	AGGGCTGAAT (GCAGCCCATT	CTCTGGAGAA	CTTCCTCACA CACCGCAGCM	120
AARGAGAAG	MCTGAAAGAC A	AAACCTGGGT	GCAGCCAGAG	AGGTCCAGAT AGATGAGCTT	180
				ACCCCAGCTG GGTCTCATTG	240
TTCCAGAACT	GCATTAGTTA A	AGATTACCCA	GACTTNGATT	TCAAAGGAAT ACTTTCATTG	300
TTCCGTCTGT	AACACGAAGT A	AATTGGGGCC	AGCTGGATGT	CAGG ATG CGT GTG GTT Met Arg Val Val -15	356
ACC ATT GT Thr Ile Va -10	A ATC TTG CTC l Ile Leu Leu	TGC TTT 1 Cys Phe C	TGC AAA GCG Cys Lys Ala	GCT GAG CTG CGC AAA Ala Glu Leu Arg Lys 1 5	404
GCA AGC CC Ala Ser Pr	A GGC AGT GTG O Gly Ser Val 10	AGA AGC (Arg Ser A	CGA GTG AAT Arg Val Asn 15	CAT GGC CGG GCG GGT His Gly Arg Ala Gly 20	452
GGA Gly					455

(2) INFORMATION FOR SEQ ID NO: 315:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 437 base pairs
 - (B) TYPE: NUCLEIC ACID
 - (C) STRANDEDNESS: DOUBLE
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: CDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: other
 - (3) LOCATION: 45..438
 - (C) IDENTIFICATION METHOD: fasta
 - (D) OTHER INFORMATION: identity 100 region 1..394

id HSU20350

(ix) FEATURE:

(A) NAME/KEY: other (B) LOCATION: 87..438

(C) IDENTIFICATION METHOD: fasta

(D) OTHER INFORMATION: identity 99 region 3..352

id HSU28934

vrt

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: 132..401

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.1

seq LLFVATLPFWTHY/LI

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 315:

AAACTCTGCA AATAAAATGC TCTTAGAGGG AAGGAAAGGG AAATACTCGT CTCTGGTAAA GTCTGAGCAG GACAGGGTGG CTGACTGGCA GATCCAGAGG TTCCCTTGGC AGTCCACGCC 120 AGGCCTTCAC C ATG GAT CAG TTC CCT GAA TCA GTG ACA GAA AAC TTT GAG 170 Met Asp Gln Phe Pro Glu Ser Val Thr Glu Asn Phe Glu -90 -85 TAC GAT GAT TTG GCT GAG GCC TGT TAT ATT GGG GAC ATC GTG GTC TTT 218 Tyr Asp Asp Leu Ala Glu Ala Cys Tyr Ile Gly Asp Ile Val Val Phe -75 GGG ACT GTG TTC CTG TCC ATA TTC TAC TCC GTC ATC TTT GCC ATT GGC 266 Gly Thr Val Phe Leu Ser Ile Phe Tyr Ser Val Ile Phe Ala Ile Gly -60 CTG GTG GGA AAT TTG TTG GTA GTG TTT GCC CTC ACC AAC AGC AAG AAG 314 Leu Val Gly Asn Leu Leu Val Val Phe Ala Leu Thr Asn Ser Lys Lys -45 -40 CCC AAG AGT GTC ACC GAC ATT TAC CTC CTG AAC CTG GCC TTG TCT GAT 362 Pro Lys Ser Val Thr Asp Ile Tyr Leu Leu Asn Leu Ala Leu Ser Asp -25 -20 -15 CTG CTG TTT GTA GCC ACT TTG CCC TTC TGG ACT CAC TAT TTG ATA AAT Leu Leu Phe Val Ala Thr Leu Pro Phe Trp Thr His Tyr Leu Ile Asn -10 GAA AAG GGC CTC CAC AAT GCC ATG TGC 437 Glu Lys Gly Leu His Asn Ala Met Cys 5

- (2) INFORMATION FOR SEQ ID NO: 316:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 35 amino acids
 - (B) TYPE: AMINO ACID

- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 11.4

seq VLALLLFVHYSNG/DE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 316:

Met Val Phe Val His Leu Tyr Leu Gly Asn Val Leu Ala Leu Leu Leu -20 -15 -10

Phe Val His Tyr Ser Asn Gly Asp Glu Ser Ser Asp Pro Gly Pro Gln -5 1 5

His Arg Ala 10

- (2) INFORMATION FOR SEQ ID NO: 317:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 34 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -29..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 11.3

seq FLLCIFLICAALA/AQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 317:

Met Gly Met Cys Phe Ala Ala Glu Ser Asp Val Gln Met Phe Ile Ala -25 -20 -15

Phe Leu Leu Cys Ile Phe Leu Ile Cys Ala Ala Leu Ala Ala Gln Lys

Ser Gly

- (2) INFORMATION FOR SEQ ID NO: 318:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 37 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -26..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 11

seq VLFLFLFWGVSLA/GS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 318:

Met Ala Val Arg Glu Leu Cys Phe Ser Arg Gln Arg Gln Val Leu Phe -25 -15

Leu Phe Leu Phe Trp Gly Val Ser Leu Ala Gly Ser Gly Phe Gly Arg -10 -5 5

Tyr Ser Val Thr Gly

- (2) INFORMATION FOR SEQ ID NO: 319:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 69 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.7 seq LILLALATGLVGG/ET

.

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 319:

Met Arg Ile Leu Gln Leu Ile Leu Leu Ala Leu Ala Thr Gly Leu Vai -15 -10 -5 Gly Gly Glu Thr Arg Ile Ile Lys Gly Phe Glu Cys Lys Pro His Ser $1 \hspace{1cm} 5 \hspace{1cm} 10$

Gln Pro Trp Gln Ala Ala Leu Phe Glu Lys Thr Arg Leu Leu Cys Gly
15 20 25 30

Ala Thr Leu Ile Ala Pro Arg Trp Leu Leu Thr Ala Ala His Cys Leu
35 40 45

Lys Pro Arg Tyr Gly 50

- (2) INFORMATION FOR SEQ ID NO: 320:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 57 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.7

seq LILLALATGLVGG/ET

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 320:
- Met Arg Ile Leu Gln Leu Ile Leu Leu Ala Leu Ala Thr Gly Leu Val
- Gly Glu Thr Arg Ile Ile Lys Gly Phe Glu Cys Lys Pro His Xaa 1 5
- Gln Pro Trp Gln Ala Ala Leu Phe Glu Lys Thr Arg Leu Leu Cys Gly 15 20 25 30

Ala Thr Leu Ile Ala Pro Arg Trp Leu
35

- (2) INFORMATION FOR SEQ ID NO: 321:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 103 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -30..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.6

seq SLLLAVLVFFLFA/LP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 321:

Met Arg Ser Cys Leu Trp Arg Cys Arg His Leu Ser Gln Gly Val Gln -30 -25 -20 -15

Trp Ser Leu Leu Ala Val Leu Val Phe Phe Leu Phe Ala Leu Pro-10 -5 1

Ser Xaa Xaa Xaa Xaa Gln Thr Lys Pro Ser Arg His Gln Arg Thr 5 10 15

Glu Asn Ile Lys Glu Arg Ser Leu Xaa Ser Leu Ala Lys Pro Lys Ser 20 25 30

Gln Ala Pro Thr Arg Ala Arg Arg Thr Thr Ile Tyr Ala Glu Pro Val 35 40 .45 50

Pro Glu Asn Asn Ala Leu Asn Thr Gln Thr Gln Pro Lys Ala His Thr
55 60 65

Thr Gly Asp Arg Arg Lys Gly
70

- (2) INFORMATION FOR SEQ ID NO: 322:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 80 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.6 seq XILLALATGLVGG/EI
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 322:

Met Arg Ile Leu Gln Xaa Ile Leu Leu Ala Leu Ala Thr Gly Leu Val

-15 -10 -5

Gly Gly Glu Ile Arg Ile Ile Lys Gly Phe Glu Cys Lys Pro His Ser $1 \hspace{1cm} 5 \hspace{1cm} 10$

Gln Pro Trp Gln Ala Ala Leu Phe Glu Lys Thr Arg Leu Leu Trp 15 20 25 30

Gly Asp Ala His Arg Pro Gln Met Ala Pro Asp Ser Ser Pro Leu Pro
35 40 45

Gln Ala Pro Leu His Ser Ser Pro Gly Ala Ala Gln Pro Pro Glu Gly
50 55 60

- (2) INFORMATION FOR SEQ ID NO: 323:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 51 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -38..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.4 seq LWLLLKLVSTXWA/VR
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 323:

Met Leu Glu Glu Cys Gly Ala Gly Val Asp Leu Gly Phe Gly Gly Val -35 -30 -25

Lys Phe Ala Ser Glu Thr Pro Asn Leu Leu Trp Leu Leu Leu Lys Leu
-20 -15 -10

Val Ser Thr Xaa Trp Ala Val Arg Val Thr Leu Ile Ile Phe Asn Asn -5 10

Gln Ala Arg

- (2) INFORMATION FOR SEQ ID NO: 324:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 27 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

PCT/IB98/01232 WO 99/06550 363

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.2

seq RCLLLALVAESSS/QT

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 324:

Met Ile Ala Cys Ser Ile Arg Glu Leu His Arg Cys Leu Leu Ala

Leu Val Ala Glu Ser Ser Ser Gln Thr His Gly

- (2) INFORMATION FOR SEQ ID NO: 325:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 32 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -17..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.2

seq SLVLCLLSATVFS/LO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 325:

Met Gly Pro Pro Ser Leu Val Leu Cys Leu Leu Ser Ala Thr Val Phe

Ser Leu Gln Gly Gly Ser Ser Ala Phe Leu Ser His His Arg Pro Gly

- (2) INFORMATION FOR SEQ ID NO: 326:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 112 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -35..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 9

seq AMWWLLLWGVLQX/XP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 326:

Met Pro Gly Pro Arg Val Trp Gly Lys Tyr Leu Trp Arg Ser Pro His -35 -25 -25

Ser Lys Gly Cys Pro Gly Ala Met Trp Trp Leu Leu Leu Trp Gly Val $-15 \qquad \qquad -10 \qquad \qquad -5$

Leu Gln Xaa Xaa Pro Asn Pro Gly Leu Arg Pro Leu Gly Xaa Arg Ala 1 5 10

Thr Pro Ala Ala Asp Ile Pro Arg Val Pro Arg Ala Val Trp Gln Arg
15 20 25

Pro Arg Glu Gln His Gly His Gln Gly Ser Arg Gly Leu Cys Cys Glu 30 40 45

Ala Arg Leu Pro Gly Leu Arg Pro Gly Ala Val Pro Gly Leu Cys Arg
50 55 60

Gly Leu Cys His Asn Leu Ile Arg Arg Phe Gly Ser Lys Pro Leu Gly
65 70 75

- (2) INFORMATION FOR SEQ ID NO: 327:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 100 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -22..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.8

sed LLTLALLGGPTWX/XK

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 327:

Met His Arg Pro Glu Ala Met Leu Leu Leu Leu Thr Leu Ala Leu Leu -20 -15 -10

Gly Gly Pro Thr Trp Xaa Xaa Lys Met Tyr Gly Pro Gly Gly Gly Lys
-5 1 5

Tyr Phe Ser Thr Thr Glu Asp Tyr Asp His Glu Ile Thr Gly Leu Arg 15 20 25

Val Ser Val Gly Xaa Leu Leu Val Lys Ser Val Gln Val Lys Leu Gly
30 35 40

Asp Ser Trp Asp Val Lys Leu Gly Gly Leu Arg Trp Glu Tyr Pro Gly
45 50 55

Ser His Pro Ala Ala Arg Arg Ile His His Lys Ser Leu Cys Arg Phe 60 65 70

Gln Ala Phe Leu 75

- (2) INFORMATION FOR SEQ ID NO: 328:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 59 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.6 seq SVSLALLSGWVGS/RQ
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 328:

Met Val Ser Val Ser Leu Ala Leu Leu Ser Gly Trp Val Gly Ser Arg

Gln Gly Gly Val Gly Leu Ser Thr Leu Val Thr Leu Gly Leu Val Ser

5 10 15

Trp Cys Trp Arg Met Val Arg Thr Gln Ala Leu Glu Gly Phe Leu Ser 20 25 30

Val Lys Tyr Tyr Ser Ala Phe Ser Ala Asp Leu 35

(2) INFORMATION FOR SEQ ID NO: 329:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 55 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -49..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.5

seq IVFLLLRVSPCLG/PS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 329:

Met His Ile Phe Ser Ile Cys Cys Met Xaa Ser Glu Leu His Lys Met -45 -40 -35

Lys Ser Leu Ser Leu Gln Leu Ala Ser Glu Lys Arg Ser Leu Val Ala
-30
-25
-20

Leu Val Glu Glu Ile Val Phe Leu Leu Leu Arg Val Ser Pro Cys Leu -15 -10 -5

Gly Pro Ser Xaa Lys Pro Arg

- (2) INFORMATION FOR SEQ ID NO: 330:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 22 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -17..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.3

seq VSALLMAWFGVLS/CV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 330:

Met Lys Leu Trp Val Ser Ala Leu Leu Met Ala Trp Phe Gly Val Leu -15 -10 -5

Ser Cys Val Gln Thr Gly

(2) INFORMATION FOR SEQ ID NO: 331:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 48 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -22..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.3

seq LLLPLMLMSMVSS/SL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 331:

Met Lys Val Leu Ile Ser Ser Leu Leu Leu Leu Leu Pro Leu Met Leu
-20 -15 -10

Met Ser Met Val Ser Ser Ser Leu Xaa Pro Gly Val Ala Arg Gly His -5 10

Arg Asp Arg Gly Gln Ala Ser Arg Arg Trp Leu Gln Glu Gly Gly Leu
15 20 25

- (2) INFORMATION FOR SEQ ID NO: 332:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 69 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -22..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.3 seq LLLPLMLMSMYSS/SL
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 332:

Met Lys Val Leu Ile Ser Ser Leu Leu Leu Leu Pro Leu Met Leu
-20 -15 -10

Met Ser Met Val Ser Ser Ser Leu Asn Pro Gly Val Ala Arg Gly His -5 1 5

Arg Asp Arg Gly Gln Ala Ser Arg Arg Trp Leu Gln Glu Gly Gln
15 20 25

Glu Cys Glu Cys Lys Asp Trp Phe Leu Arg Ala Pro Arg Arg Lys Phe 30 35 40

Met Thr Val Ser Gly 45

- (2) INFORMATION FOR SEQ ID NO: 333:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 16 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -14..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.2 seq LLLLQLSLPSPTS/SP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 333:

Met Leu Leu Leu Gln Leu Ser Leu Pro Ser Pro Thr Ser Ser Pro -10 -5 1

- (2) INFORMATION FOR SEQ ID NO: 334:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 27 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:

- (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -17..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.1 seq LSFKLLLLAVALG/FF
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 334:

Met Leu Lys Met Leu Ser Phe Lys Leu Leu Leu Leu Ala Val Ala Leu
-15 -5

Gly Phe Phe Glu Gly Asp Ala Lys Phe Gly Glu
1 5 10

- (2) INFORMATION FOR SEQ ID NO: 335:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 69 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -22..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8

seq LLTLALLGXXXWA/GK

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 335:
- Met His Arg Pro Glu Ala Met Leu Leu Leu Leu Thr Leu Ala Leu Leu -20 -15 -10
- Gly Xaa Xaa Xaa Trp Ala Gly Lys Met Tyr Gly Pro Gly Gly Gly Lys
 -5 5 10
- Tyr Phe Ser Thr Thr Glu Asp Tyr Asp His Glu Ile Thr Gly Leu Arg
- Val Ser Val Gly Leu Leu Leu Val Lys Ser Val Gln Val Lys Leu Gly 30 35 40

Asp Ser Trp Asp Val 45

- (2) INFORMATION FOR SEQ ID NO: 336:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 70 amino acids

- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -16..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8

seq VSAVLCVCAAAWC/SQ

PCT/IB98/01232

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 336:

Met Leu Lys Val Ser Ala Val Leu Cys Val Cys Ala Ala Ala Trp Cys
-15 -5

Ser Gln Ser Leu Ala Ala Ala Ala Ala Val Ala Ala Ala Gly Gly Arg 1 5 10 15

Ser Asp Gly Gly Asn Phe Leu Asp Asp Lys Gln Trp Leu Thr Thr Ile 20 25 30

Ser Gln Tyr Asp Lys Glu Val Gly Gln Trp Asn Lys Phe Arg Asp Asp 35 40 45

Asp Tyr Phe Arg Thr Gly 50

- (2) INFORMATION FOR SEQ ID NO: 337:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 45 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -17..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.8 seq VLWLISFFTFTDG/HG
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 337:

Met Lys Val Gly Val Leu Trp Leu Ile Ser Phe Phe Thr Phe Thr Asp
-15 -10 -5

Gly His Gly Gly Phe Leu Gly Lys Asn Asp Gly Ile Lys Thr Lys Lys
1 5 10 15

Glu Leu Ile Val Asn Lys Lys His Leu Gly Leu Gly
20 25

- (2) INFORMATION FOR SEQ ID NO: 338:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 19 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -16..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.7 seq ILLDLICLLFITA/CV
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 338:

Met Cys Ile Ile Leu Leu Asp Leu Ile Cys Leu Leu Phe Ile Thr Ala
-15 -10 -5

Cys Val Gly

- (2) INFORMATION FOR SEQ ID NO: 339:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 62 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -59..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.6 seq FMVFGSFFPLISC/OP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 339:

Met Asp Cys Ala Ser Ile Ser Val Lys Phe Thr Ser Met Ala Thr Met
-55 -50 -45

His Asp Leu Ser Gln Phe Trp Ala Ser Arg Gly Glu Val Thr Asn Trp
-40 -35 -30

Trp Pro Val Gly Gln Thr Ser Leu Pro Leu Phe Tyr Leu Ala Phe Met
-25 -20 -15

Val Phe Gly Ser Phe Phe Pro Leu Ile Ser Cys Gln Pro Gly
-10 -5 1

- (2) INFORMATION FOR SEQ ID NO: 340:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 57 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -20..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.6 seq LVVLFGITAGATG/AK
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 340:

Met Thr Ala Ser Pro Asp Tyr Leu Val Val Leu Phe Gly Ile Thr Ala -20 -15 -10 -5

Gly Ala Thr Gly Ala Lys Leu Gly Ser Asp Glu Lys Glu Leu Ile Leu 1 5 10

Leu Phe Trp Lys Val Val Asp Leu Ala Asn Lys Lys Val Gly Gln Leu
15 20 25

His Glu Xaa Xaa Leu Asp Arg Ile Trp 30 35

- (2) INFORMATION FOR SEQ ID NO: 341:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 79 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.6

seq CVLVLAAAAGAVA/VF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 341:

Met Val Cys Val Leu Val Leu Ala Ala Ala Ala Gly Ala Val Ala Val -15 -5 1

Phe Leu Ile Leu Arg Ile Trp Val Val Leu Arg Ser Met Asp Val Thr 5 10 15

Pro Arg Glu Ser Leu Ser Ile Leu Val Val Ala Gly Ser Gly Gly His 20 25 30

Thr Thr Glu Ile Leu Arg Leu Leu Gly Ser Leu Ser Asn Ala Tyr Ser 35 40 45

Pro Arg His Tyr Val Ile Ala Asp Thr Asp Glu Met Ser Ala Thr 50 55 60

- (2) INFORMATION FOR SEQ ID NO: 342:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 82 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -44..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.5 seq LMIPLLLLTPITA/TS
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 342:

Met Lys Lys Thr Gly Asp Gly Gly Thr Leu Ser Thr Glu Arg Ile Gly
-40 -35 -30

Gly Ala Ala Leu Leu Ser Leu Leu Leu Lys Arg Met Lys Met Thr Leu
-25 -20 -15

Met Ile Pro Leu Leu Leu Thr Pro Ile Thr Ala Thr Ser Thr Ser -10 -5 1

Arg Trp Pro Glu Ile Gly Val Val Ala Ile Arg Ser Gln Leu Arg Ala 5 10 15 20

Leu His Thr Cys Gly Gln Glu Pro Val Pro Ala Met Gly Ser Glu Gly 25 30 35

Ala Ala

- (2) INFORMATION FOR SEQ ID NO: 343:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 103 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.5 seq LTFLQLLLISSLP/RE
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 343:

Met Glu Leu Gly Cys Trp Thr Gln Leu Gly Leu Thr Phe Leu Gln Leu
-20 -15 -10

Leu Leu Ile Ser Ser Leu Pro Arg Glu Tyr Thr Val Ile Asn Glu Ala -5 1 5

Cys Pro Gly Ala Glu Trp Xaa Ile Met Cys Arg Glu Cys Cys Glu Tyr 10 15 20 25

Asp Gln Ile Glu Cys Val Cys Pro Gly Lys Arg Glu Val Val Gly Tyr 30 35

Thr Ile Pro Cys Cys Arg Asn Glu Xaa Asn Glu Cys Asp Ser Cys Leu
45 50 55

Ile His Pro Gly Cys Thr Ile Phe Glu Asn Cys Xaa Ser Cys Arg Asn 60 65 70

Gly Ser Trp Gly Gly Thr Leu 75 80

(2) INFORMATION FOR SEQ ID NO: 344:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 80 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -27..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.2

seq SLLFFLLLEGGXT/EQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 344:

Met Arg Xaa Lys Trp Lys Met Gly Gly Met Lys Tyr Ile Phe Ser Leu
-25 -20 -15

Leu Phe Phe Leu Leu Glu Gly Gly Xaa Thr Glu Gln Val Xaa His -10 -5 1 5

Ser Glu Thr Tyr Cys Met Phe Gln Asp Lys Lys Tyr Arg Val Gly Glu 10 15 20

Arg Trp His Pro Tyr Leu Glu Pro Tyr Gly Leu Val Tyr Cys Val Asn 25 30

Cys Ile Cys Ser Glu Xaa Gly Asn Val Leu Cys Ser Arg Val Arg Cys 40 45 50

- (2) INFORMATION FOR SEQ ID NO: 345:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 81 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -19..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.2

seg VSIMLLLVTVSDC/AV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 345:

Met Arg Gly Ala Thr Arg Val Ser Ile Met Leu Leu Leu Val Thr Val -15 -10 -5

Ser Asp Cys Ala Val Ile Thr Gly Ala Cys Glu Arg Asp Val Gln Cys
l 5 10

Gly Ala Gly Thr Cys Cys Ala Ile Ser Leu Trp Leu Arg Gly Leu Arg 15 20 25

Met Cys Thr Pro Leu Gly Arg Glu Gly Glu Glu Cys His Pro Gly Ser 30 40 45

His Lys Ile Pro Phe Phe Arg Lys Arg Lys His His Thr Cys Pro Cys 50 55 60

Leu

- (2) INFORMATION FOR SEQ ID NO: 346:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 47 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.2 seq SALLFSLLCEAST/VV
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 346:

Met Ile Ala Ile Ser Ala Val Ser Ser Ala Leu Leu Phe Ser Leu Leu -20 -15 -10

Cys Glu Ala Ser Thr Val Val Leu Leu Asn Ser Thr Asp Ser Ser Pro
-5 1 5 10

Xaa Thr Asn Asn Phe Xaa Asp Xaa Glu Ala Ala Leu Lys Ala His

- (2) INFORMATION FOR SEQ ID NO: 347:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 85 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

377

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.2

seq SALLFSLLCEAST/VV

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 347:
- Met Ile Ala Ile Ser Ala Val Ser Ser Ala Leu Leu Phe Ser Leu Leu
- Cys Glu Ala Ser Thr Val Val Leu Leu Asn Ser Thr Asp Ser Ser Pro
- Pro Thr Asn Asn Phe Thr Asp Ile Glu Ala Ala Leu Lys Ala Gln Leu
- Asp Ser Ala Asp Ile Pro Lys Ala Arg Arg Lys Arg Tyr Ile Ser Gln
- Asn Asp Met Ile Ala Ile Leu Asp Tyr His Asn Gln Val Arg Gly Lys

Val Phe Pro Xaa Ala 60

- (2) INFORMATION FOR SEQ ID NO: 348:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 25 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -22..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.2

seq LLTLVLCVAVAYE/RQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 348:

Met Asp Pro Ash Gly Gly Cys Cys Thr Leu Leu Thr Leu Val Leu Cys -15

PCT/IB98/01232

Val Ala Val Ala Tyr Glu Arg Gln Glu -5

- (2) INFORMATION FOR SEQ ID NO: 349:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 34 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -25..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.2

seq LFTFSTSLPSSLS/SS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 349:

Met Glu Gly Glu Ile Tyr Phe Gln Val Phe Leu Ser Leu Phe Thr Phe -25 -20 -15 -10

Ser Thr Ser Leu Pro Ser Ser Leu Ser Ser Ser Ser Leu Ser Ser Ser Ser 5

Asn Gly

- (2) INFORMATION FOR SEQ ID NO: 350:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 45 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -41..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7

seq FLCMLAAIDLALS/TS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 350:

Met Tyr Val Val Ala Met Phe Gly Asn Cys Ile Val Val Phe Ile Val -40 -35 -30

Arg Thr Glu Arg Ser Leu His Ala Pro Met Tyr Leu Phe Leu Cys Met -25 -15 -10

Leu Ala Ala Ile Asp Leu Ala Leu Ser Thr Ser Thr Met

- (2) INFORMATION FOR SEQ ID NO: 351:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 63 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -43..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7

seq PWFLAPWCPGTQS/NR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 351:

Met Arg Glu Thr Xaa Pro Leu Pro Lys Pro Leu Lys Asp Thr Ala Pro
-40 -35 -30

Ser Ser His Gly Val Gly Ser Asp Ser Pro Ser Ala Thr Arg Pro Trp -25 -15

Phe Leu Ala Pro Trp Cys Pro Gly Thr Gln Ser Asn Arg Ile Cys His

Pro Pro Leu Ser Ser Pro Pro Asp Gln Ala Thr Cys Leu Arg Gly 10 15 20

- (2) INFORMATION FOR SEQ ID NO: 352:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 93 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate

WO 99/06550 PCT/IB98/01232

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -60..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7 seq VLVVLALRSLGRS/CS
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 352:

Met Asp Arg Pro Gly Ser Leu Ser Val Phe Gly Ser Leu Pro Ala Ser -60 -55 -50 -45

Leu Gly Thr Trp Leu Ser Ser Pro Ala Trp Leu Val Asp Arg Pro Val -40 -35 -30

Arg Ser Ala His Pro Ser Ala Asn Ser Thr Gly Val Arg Met Ser Val -25 -20 -15

Leu Val Val Leu Ala Leu Arg Ser Leu Gly Arg Ser Cys Ser Leu Ser -10 -5 1

Gln Ala Ala Pro Ser Arg Trp Thr Arg Ser Asn Asp Ala Pro Gln Pro 5 10 15 20

Pro Gly Ser Gln His Ile Phe His Thr Xaa Val Pro Gly 25 30

- (2) INFORMATION FOR SEQ ID NO: 353:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 36 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7

seq VILLFSYPSCCLC/FL

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 353:
- Met His Tyr Phe Val Ala Gly Lys Val Ile Leu Leu Phe Ser Tyr Pro -20 -15 -10

Ser Cys Cys Leu Cys Phe Leu Val Tyr Arg Arg Val Ser Xaa Leu Phe $^{-5}$ 1 5 10

Lys Cys Phe Glu

15

WO 99/06550 PC

(2) INFORMATION FOR SEO ID NO: 354:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 53 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -19..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7

seq STVVLQVLTQATS/QD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 354:

Met Asp Leu Asn Ser Ala Ser Thr Val Val Leu Gln Val Leu Thr Gln
-15 -10 -5

Ala Thr Ser Gln Asp Thr Ala Val Leu Lys Pro Ala Glu Glu Gln Leu
1 5 10

Lys Gln Trp Glu Thr Gln Pro Gly Phe Tyr Ser Val Leu Leu Asn Ile 15 20 25

Phe Thr Asn His Gly 30

- (2) INFORMATION FOR SEQ ID NO: 355:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 77 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -73..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7

seg FLCMLAAIDLALS/TS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 355:

Met Ser Ser Cys Asn Phe Thr His Ala Thr Phe Val Leu Ile Gly Ile - -70 -65 -60

Pro Gly Leu Glu Lys Ala His Phe Trp Val Gly Phe Pro Leu Leu Ser

Met Tyr Val Val Ala Met Phe Gly Asn Cys Ile Val Val Phe Ile Val -40 -35 -30

Arg Thr Glu Arg Ser Leu His Ala Pro Met Tyr Leu Phe Leu Cys Met -25 -15 -10

Leu Ala Ala Ile Asp Leu Ala Leu Ser Thr Ser Thr Met -5

(2) INFORMATION FOR SEQ ID NO: 356:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 79 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -56..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.9 seq PLFFSCSISATHS/CV
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 356:

Met Tyr Arg Leu Ser Leu Ile Ala Gly Pro Gly Ser Tyr Pro Val Leu
-55 -50 -45

Arg Trp Gly Val Trp Asp Ile Pro Ser Ser Leu Val Gln Val Thr Tyr -40 -35 -30 -25

His Gln Pro Asn Leu Thr Thr Asn Leu Asp Leu Pro Leu Phe Phe Ser

Cys Ser Ile Ser Ala Thr His Ser Cys Val Lys Pro Pro Ser Val Ile -5

Ile Gly Ile Ser Ser Phe Leu Ser Phe Pro Tyr Gln Thr Leu Val 10 15 20

(2) INFORMATION FOR SEQ ID NO: 357:

WO 99/06550 383

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 91 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -24..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.9

seq LCFLLLAVAMSFF/GS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 357:

Met Leu Val Asp Gly Pro Ser Glu Arg Pro Ala Leu Cys Phe Leu Leu -20 -15 -10

Leu Ala Val Ala Met Ser Phe Phe Gly Ser Ala Leu Ser Ile Asp Glu -5 1 5

Thr Arg Ala His Leu Leu Lys Glu Lys Met Met Arg Leu Gly Gly
10 15 20

Arg Leu Val Leu Asn Thr Lys Glu Glu Leu Ala Asn Glu Arg Leu Met 30 35 40

Thr Leu Lys Ile Ala Glu Met Lys Glu Ala Met Arg Thr Leu Ile Phe \$45\$ 50 55

Pro Pro Ser Met His Phe Phe Gln Ala Lys Trp
60 65

- (2) INFORMATION FOR SEQ ID NO: 358:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 60 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -35..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.9

seq XLXXLLTPPPSYG/HQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 358:

Met Pro Cys Ser Leu Thr Trp Arg Leu Pro Pro Arg Thr Cys Gln Xaa -35 -25 -20

Xaa Gly Leu Xaa Lys Ser Xaa Leu Xaa Xaa Leu Leu Thr Pro Pro Pro -15 -10 -5

Ser Gly Asp Lys Asp His Leu Tyr Ser Thr Val Cys
15 20 25

- (2) INFORMATION FOR SEQ ID NO: 359:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 85 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -41..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.8 seq LFLFLTSIAEXCS/TP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 359:
- Met Val Xaa Trp Leu Val Leu Phe Ala Leu Gln Ile Tyr Ser Tyr Xaa-40 -35 -30
- Ser Thr Arg Asp Gln Pro Ala Ser Arg Xaa Arg Leu Leu Phe Leu Phe -25 -20 -15 -10

Leu Thr Ser I-le Ala Glu Xaa Cys Ser Thr Pro Tyr Ser Leu Leu Gly

Xaa Val Phe Thr Val Ser Phe Val Ala Leu Gly Val Leu Thr Leu Cys
10 20

Lys Phe Tyr Leu Gln Gly Tyr Arg Ala Phe Met Asn Asp Pro Ala Met 25 30 35

Asn Arg Gly Gly Ala

(2) INFORMATION FOR SEQ ID NO: 360:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 20 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.7

seq LPLLXXXSLPVGA/WL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 360:

Met Ala Arg His Gly Leu Pro Leu Leu Xaa Xaa Xaa Ser Leu Pro Val

Gly Ala Trp Leu 1

- (2) INFORMATION FOR SEQ ID NO: 361:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 50 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -37..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.7

seq ILYILWYCSVCSS/GS

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 361:
- Met Val His Leu Arg Thr Gly Leu Met Leu Met Ser Ala Asp Arg Leu
 -35 -30 -25
- Arg Thr Leu Tyr Tyr Thr Val Thr Ile Leu Tyr Ile Leu Trp Tyr Cys
 -20 -15 -10
- Ser Val Cys Ser Ser Gly Ser Leu Leu Ser Thr Ser Ile Met Lys Lys -5 1 5 10

Arg Met

- (2) INFORMATION FOR SEQ ID NO: 362:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 51 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.7

seq ILSTVTALTFARA/LD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 362:

Met Gly Ile Leu Ser Thr Val Thr Ala Leu Thr Phe Ala Arg Ala Leu
-15 -5 1

Asp Gly Cys Arg Asn Gly Ile Ala His Pro Ala Ser Glu Lys His Arg 5 10 15

Leu Glu Lys Cys Arg Glu Leu Glu Ser Ser His Ser Ala Pro Gly Ser 20 25 30

Thr Gln Gln 35

- (2) INFORMATION FOR SEQ ID NO: 363:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 34 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.5 seq LTFLQXLLISSLX/RE

PCT/IB98/01232 WO 99/06550 387

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 363:

Met Glu Leu Gly Cys Trp Thr Gln Leu Gly Leu Thr Phe Leu Gln Xaa -20 -15

Leu Leu Ile Ser Ser Leu Xaa Arg Glu Tyr Thr Val Ile Asn Glu Ala

Arg Lys 10

- (2) INFORMATION FOR SEQ ID NO: 364:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 36 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -22..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.4 seq FLLCXSVFTDCKG/DV
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 364:

Met Glu Leu Leu Arg Val Cys Ser Phe Phe Leu Leu Cys Xaa Ser Val -15

Phe Thr Asp Cys Lys Gly Asp Val Leu Cys Val Lys Met Glu Gln Ser

Gln Ile Cys Ala

- (2) INFORMATION FOR SEQ ID NO: 365:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 29 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate

WO 99/06550 PCT/IB98/01232

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -22..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.3 seq TWFLLLPPGQCRA/VG
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 365:

Met Ile Val Arg Pro Arg Leu Asn Leu Thr Trp Phe Leu Leu Pro -20 -15 -10

Pro Gly Gln Cys Arg Ala Val Gly Ala Thr Trp Pro Gly
-5 1 5

- (2) INFORMATION FOR SEQ ID NO: 366:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 40 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -19..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.3 seq MVALCCCLWKISG/CE
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 366:

Met Gln Phe Leu Phe Lys Met Val Ala Leu Cys Cys Cys Leu Trp Lys

Ile Ser Gly Cys Glu Glu Val Pro Leu Thr Tyr Asn Leu Leu Lys Cys

1 5 10

Leu Leu Asp Lys Ala His Val Gly
15 20

- (2) INFORMATION FOR SEQ ID NO: 367:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 66 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

WO 99/06550 PCT/IB98/01232

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.3

seq CVCAAAXXSQSLX/XX

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 367:

Met Leu Lys Val Ser Ala Val Leu Cys Val Cys Ala Ala Ala Xaa Xaa -20 -15 -10

Ser Gln Ser Leu Xaa Xaa Xaa Ala Ala Val Ala Ala Ala Gly Gly Arg -5 1 10

Ser Asp Gly Gly Asn Phe Leu Asp Asp Lys Gln Trp Leu Thr Xaa Ile 15 20 25

Ser Gln Tyr Asp Lys Glu Xaa Gly Xaa Trp Asn Lys Phe Arg Asp Asp 30 35 40

Xaa Tyr 45

- (2) INFORMATION FOR SEQ ID NO: 368:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 57 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.3

seq MVALCCCLWKISG/CE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 368:

Met Ser Met Gln Phe Leu Phe Lys Met Val Ala Leu Cys Cys Cys Leu
-20 -15 -10

Trp Lys Ile Ser Gly Cys Glu Glu Val Pro Leu Thr Tyr Asn Leu Leu -5 1 5 10

Lys Cys Leu Leu Asp Lys Ala His Cys Val Leu Leu Thr Pro Cys Gly

15 20 25

Tyr Ile Phe Ser Leu Ile Ser Pro Gly 30 35

- (2) INFORMATION FOR SEQ ID NO: 369:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -17..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.2

seq LWILLGSLSCRTS/NR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 369:

Met Ala Gln His Leu Trp Ile Leu Leu Gly Ser Leu Ser Cys Arg Thr
-15 -10 -5

Ser Asn Arg Arg

- (2) INFORMATION FOR SEQ ID NO: 370:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 59 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -28..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.1 seq LYLFSGFWTFXLG/KF
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 370:

Met Asn Lys Glu Xaa Val Ser Xaa Glu Arg Xaa Ala Gln Val Arg Leu
-25 -20 -15

Tyr Leu Phe Ser Gly Phe Trp Thr Phe Xaa Leu Gly Lys Phe Lys Gln -10 -5 1

Gly Glu Xaa Ser Tyr Xaa Xaa Ile Leu Glu Arg Leu Leu Trp Gln Gln 5 10 15 20

Gln Tyr Xaa Gly Trp Leu Val Gly Asp Lys Arg
25 30

- (2) INFORMATION FOR SEQ ID NO: 371:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 80 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -54..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6

seq IVFIFLILLNTAA/QV

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 371:
- Met Val Leu Trp Arg Ala Lys Ile Xaa Arg Asn Val Pro Val Thr Leu -50 -45 -40
- Ser Glu Glu Asn Arg Ser Glu Gly Lys Val Gly Phe Gln Ala Tyr Lys -35 -30 -25
- Asn Tyr Phe Arg Ala Gly Ala His Trp Ile Val Phe Ile Phe Leu Ile
 -20 -15 -10
- Leu Leu Asn Thr Ala Ala Gln Val Ala Tyr Val Leu Gln Asp Trp Trp
 -5 1 5
- Leu Ser Tyr Trp Ala Asn Lys Gln Ser Met Leu Asn Val Thr Val Asn 15 20 25
- (2) INFORMATION FOR SEQ ID NO: 372:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 36 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR

- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6

seq FTSVLWLTSPSQP/NT

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 372:

Met Leu Leu Xaa Phe Phe Thr Ser Val Leu Trp Leu Thr Ser Pro Ser -15 -10 -5

Gln Pro Asn Thr Cys Pro Ser Ser Leu Leu Cys Thr Tyr Pro Asn Leu

1 5 10

Asn Pro Pro Trp 15

- (2) INFORMATION FOR SEQ ID NO: 373:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 30 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -22..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq IILGCLALFLLLO/RK

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 373:
- Met Glu Leu Ile Ser Pro Thr Val Ile Ile Ile Leu Gly Cys Leu Ala -20 -15 -10
- Leu Phe Leu Leu Gln Arg Lys Asn Leu Arg Arg Pro Trp
 -5 5
- (2) INFORMATION FOR SEQ ID NO: 374:
 - (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 53 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -47..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq TWLGLLSFQNLHC/FP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 374:

Met His Gly Phe Glu Ile Ile Ser Leu Lys Glu Glu Ser Pro Leu Gly
-45 -35

Lys Val Ser Gln Gly Pro Leu Phe Asn Val Thr Ser Gly Ser Ser Ser -30 -25 -20

Pro Val Thr Trp Leu Gly Leu Leu Ser Phe Gln Asn Leu His Cys Phe -15 -10 -5 1

Pro Asp Leu Pro Gly

- (2) INFORMATION FOR SEQ ID NO: 375:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 63 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (-ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -56..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq NTLFLHLSGLSAA/DT

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 375:

Met Thr Trp Val Arg His Ala Pro Gly Lys Ser Leu Glu Trp Val Ala
-55 -45

Thr Val Thr Asp Gly Gly Asp Lys Thr Phe Tyr Ala Ala Ser Val Lys

WO 99/06550 PCT/IB98/01232

-40 -35

-30

25

Gly Arg Phe Asn Val Ser Arg Asp Asn Ser Lys Asn Thr Leu Phe Leu -20 -15 -10

His Leu Ser Gly Leu Ser Ala Ala Asp Thr Gly Trp Trp Gly Ile
-5 1 5

- (2) INFORMATION FOR SEQ ID NO: 376:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 28 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -14..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.8

seq LTSFFSLTANCQS/AG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 376:

Met Leu Thr Ser Phe Phe Ser Leu Thr Ala Asn Cys Gln Ser Ala Gly -10 -5 1

Thr Ile Ser Phe Ala Ala Phe Ser Leu Met Pro Gly 5 10

- (2) INFORMATION FOR SEQ ID NO: 377:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (3) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.8 seq LTPLFFMXPTGFS/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 377:

Met Leu Cys Leu Leu Thr Pro Leu Phe Phe Met Xaa Pro Thr Gly
-15 -10 -5

Phe Ser Ser Pro Ser Pro Gly

- (2) INFORMATION FOR SEQ ID NO: 378:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 37 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.7 seq HSLFLSLLGLCPS/KT
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 378:
- Met Asp Asp Asp Tyr Glu Ala Tyr His Ser Leu Phe Leu Ser Leu Leu -20 -15 -10 .
- Gly Leu Cys Pro Ser Lys Thr Pro Ile Asn Glu Asn Ala Pro Val Phe -5 1 5 10

Asp Pro Glu Pro Val

- (2) INFORMATION FOR SEQ ID NO: 379:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 32 amino acids
 - (3) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -19..-1

PCT/IB98/01232 396

- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 5.7 seq WLVWLLLGHMVVS/QM

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 379:

Met Glu Trp Gly Lys Gln Trp Leu Val Trp Leu Leu Leu Gly His Met -10

Val Val Ser Gln Met Ala Thr Leu Leu Ala Arg Lys His Arg Pro Trp

- (2) INFORMATION FOR SEQ ID NO: 380:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 49 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -39..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.7 seq LTQGVLWILVIQA/VP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 380:

Met Arg Arg Gly Lys Arg Leu Leu Glu Ser Gln Ser Ser Ser Pro Lys

Ala Cys Leu Gln Leu Gly Phe Glu Thr Glu Leu Thr Gln Gly Val Leu

Trp Ile Leu Val Ile Gln Ala Val Pro Val Pro Ser Leu Thr Lys Thr

Lys 10

- (2) INFORMATION FOR SEQ ID NO: 381:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 24 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -20..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.7

seq ALLESVVWLPCHG/RG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 381:

Met Val Ala Ala Thr Glu Ala Ala Leu Leu Glu Ser Val Val Trp Leu -20 -15 -10 -5

Pro Cys His Gly Arg Gly Gly Ser

- (2) INFORMATION FOR SEQ ID NO: 382:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 22 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -19..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.6

seq VSLPLLSSWGSTA/WT

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 382:

Met Ser Trp Asn Pro Ser Val Ser Leu Pro Leu Leu Ser Ser Trp Gly
-15 -10 -5

Ser Thr Ala Trp Thr Leu 1

- (2) INFORMATION FOR SEQ ID NO: 383:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 47 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Prostate

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: -22..-1

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 5.6

seq LILLSLHLERRWT/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 383:

Met Lys Arg Ile Gln Gly Ile Leu Phe Leu Ile Leu Ser Leu His
-20 -15 -10

Leu Glu Arg Arg Trp Thr Ser Pro Ser Asp His Ser Leu Leu Gly
-5 1 5 10

Gly Asn Ser Leu Ala Gln His Ala Glu Ser Val Val Arg Gln Gly
15 20 25

- (2) INFORMATION FOR SEQ ID NO: 384:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 46 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -35..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.5

seq LLTFGLEVCLAAG/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 384:

Met Val Gln Arg Leu Trp Val Ser Arg Leu Leu Arg His Arg Lys Ala
-35 -30 -25 -20

Gln Leu Xaa Leu Xaa Asn Leu Leu Thr Phe Gly Leu Glu Val Cys Leu -15 -10 -5

Ala Ala Gly Ser Pro Met Cys Arg Leu Cys Cys Trp Lys Trp

1 5 10

(2) INFORMATION FOR SEQ ID NO: 385:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 122 amino acids

(B) TYPE: AMINO ACID

(D) TOPOLOGY: LINEAR

- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.5 seq PFALVTSCSSVFS/GD
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 385:

Met Ala Ala Gly Val Pro Phe Ala Leu Val Thr Ser Cys Ser Ser Val -15 -10 -5

Phe Ser Gly Asp Gln Leu Val Gln His Ile Leu Gly Thr Glu Asp Leu
1 5 10

Ile Val Glu Val Thr Ser Asn Asp Ala Val Arg Phe Tyr Pro Trp Thr 15 20 \cdot 25 30

Ile Asp Asn Lys Tyr Tyr Ser Ala Asp Ile Asn Leu Cys Val Val Pro $35 \hspace{1cm} 40 \hspace{1cm} 45$

Asn Lys Phe Leu Val Thr Ala Glu Ile Ala Glu Ser Val Gln Ala Phe
50 55 60

Val Val Tyr Phe Asp Xaa Thr Gln Xaa Ser Gly Leu Asp Ser Val Ser 65 70 75

Ser Trp Leu Pro Leu Ala Lys Ala Trp Leu Pro Glu Val Met Ile Leu 89 85 90

Val Cys Asp Arg Val Ser Glu Asp Gly Ile 95 . 100

- (2) INFORMATION FOR SEQ ID NO: 386:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 amino acids
 - (3) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -14..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.5

seq TVFLXFCFPRCHS/DS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 386:

Met Thr Val Phe Leu Xaa Phe Cys Phe Pro Arg Cys His Ser Asp Ser -10 -5 1

His Xaa Xaa Gln Gln Ser Ala

- (2) INFORMATION FOR SEQ ID NO: 387:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 89 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -48..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.4

seq ILLEVFVWNGLQG/LP

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 387:
- Met Xaa Pro Asn Asn Phe Trp Gln Lys Leu Gly Arg Lys Lys Pro Arg
 -45 -40 -35
- Ile Phe Thr Cys Thr Gln Ser Ser Thr Gly Glu Ala Ala Val Lys Ala -30 -25 -20
- Glu Asn Leu Ile Leu Leu Glu Val Phe Val Trp Asn Gly Leu Gln Gly -15 -5
- Leu Pro Ser Glu Leu Ser Asp Thr Ser Gly Ser Ser Lys Lys Leu Gly
 1 5 10 15
- Ser Leu Val Gly Trp Trp Arg Thr Leu Lys Met Ala Pro Ala Cys Leu 20 25 30
- Trp Ser Met Trp Glu Ser Pro Pro Arg
 35

- (2) INFORMATION FOR SEQ ID NO: 388:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 73 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -36..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.3 seq ALYIMCVPHSVWG/CA
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 388:

Met Phe Arg Ser Asp Arg Met Trp Xaa Cys His Trp Lys Trp Lys Pro
-35 -30 -25

Ser Pro Leu Leu Phe Leu Phe Ala Leu Tyr Ile Met Cys Val Pro His -20 -15 -10 -5

Ser Val Trp Gly Cys Ala Asn Cys Arg Val Val Leu Ser Asn Pro Ser 1 5 10

Gly Thr Phe Thr Ser Pro Cys Tyr Pro Asn Asp Tyr Pro Asn Ser Gln
15 20 25

Ala Cys Met Trp Thr Leu Arg Asp Pro

- (2) INFORMATION FOR SEQ ID NO: 389:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 92 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -31..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.3 seq LVALSSELPFLGA/GV
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 389:

Met Thr Gln Arg Ser Ile Ala Gly Pro Ile Cys Asn Leu Lys Phe Val -30 -25 -20

Thr Leu Leu Val Ala Leu Ser Ser Glu Leu Pro Phe Leu Gly Ala Gly
-15 -5 1

Val Gln Leu Gln Asp Asn Gly Tyr Asn Gly Leu Leu Ile Ala Ile Asn 5 10 15

Pro Gln Val Pro Glu Asn Gln Asn Leu Ile Ser Asn Ile Lys Glu Met
20 25 30

Phe Phe Arg Asn Ile Lys Ile Leu Ile Pro Ala Gln 50 55 60

(2) INFORMATION FOR SEQ ID NO: 390:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 46 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
- (iz) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -14..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.3

seq IIPLLLLRSACN/VH

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 390:

Met Ile Ile Pro Leu Leu Leu Leu Arg Ser Ala Cys Asn Val His
-10 -5 1

Leu Pro His Gln Thr Ala Ser Pro Ala Ser Leu Ser Pro Gln Gly Leu
5 10 15

Ala Trp Gly Leu Leu His Gly Gly Cys Ser Val Thr Val Arg
20 25 30

(2" INFORMATION FOR SEQ ID NO: 391:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 21 amino acids
 - (B) TYPE: AMINO ACID

- .(D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -19..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.3

seq VLLLSXNLNLIIQ/SS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 391:

Met Xaa Ser Pro Leu Pro Val Leu Leu Ser Xaa Asn Leu Asn Leu -15 -10 -5

Ile Ile Gln Ser Ser

- (2) INFORMATION FOR SEQ ID NO: 392:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 53 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -46..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.2

seq LLTFLVFTXKLSS/LN

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 392:

Met Leu Met Cys Lys Met Leu Lys Ser Gln Lys Asn Cys Gln Glu Asn -45 -35

Xaa Xaa Ile Lys Ile Ile Leu Phe Leu Lys Pro Met Cys Ser Pro Gln
-30 -25 -20 -15

Tyr Leu Leu Thr Phe Leu Val Phe Thr Xaa Lys Leu Ser Ser Leu Asn -10 -5 1

Ile Kaa Lys Phe His

- (2) INFORMATION FOR SEQ ID NO: 393:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 55 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -52..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.2 seq IIVILHCAASIIS/CP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 393:
- Met Lys Lys Ser Ser Pro Asn Gln Tyr Leu His Ser Ser Leu His -50 -45 -40
- Xaa Ile Arg Leu Phe Ser Phe Leu His Phe Ser Glu Glu Gly Val Leu
 -35 -30 -25
- Leu Leu Ala Ile Asp Leu Lys Ile Ile Val Ile Leu His Cys Ala Ala -20 -15 -10 -5

Ser Ile Ile Ser Cys Pro Ser

- (2) INFORMATION FOR SEO ID NO: 394:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 73 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.1

seq ATSVSLEAQSCFA/WP

(zi) SEQUENCE DESCRIPTION: SEQ ID NO: 394:

Met Phe Ser Cys Phe Phe Ser Thr Ser Leu Ala Thr Ser Val Ser Leu
-20 -15 -10

Glu Ala Gln Ser Cys Phe Ala Trp Pro Leu Ile Val Ser Phe Pro Gln
-5 5

Gly Ser Leu Leu Ser Pro Phe Leu Leu Met Ser Tyr Asn Leu Ser His 10 20 25

Leu Ile Tyr Ser Gly Glu Leu Asn Gly Arg Leu Tyr Ala Glu Asn Ser 30 35 40

Gln Ile Cys Ile Cys Ser Pro Ala Gly
45 50

(2) INFORMATION FOR SEQ ID NO: 395:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 75 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -50..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.1

seq RTALILAVCCGSA/SI

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 395:

Het His-His Gly Leu Thr Pro Leu Leu Gly Val His Glu Gln Lys
-50 -45 -40 -35

Heu Asp Arg Tyr Gly Arg Thr Ala Leu Ile Leu Ala Val Cys Cys Gly
-15 -10 -5

Ser Ala Ser Ile Val Ser Leu Leu Glu Gln Asn Ile Asp Val Ser 1 5 10

Ser Gin Asp Leu Ser Gly Gln Thr Ala Pro Gly 15 20 25

- (2) INFORMATION FOR SEQ ID NO: 396:
 - (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 21 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -17..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.1

seq IYFFACFQALTSS/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 396:

Met Ser Pro Cys Ile Tyr Phe Phe Ala Cys Phe Gln Ala Leu Thr Ser -15 -10 -5

Ser Ser Pro Pro Gln
1

- (2) INFORMATION FOR SEQ ID NO: 397:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 90 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -31..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.1

seq VSGASGFLPPARS/RI

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 397:
- Met Ala Glu Glu Met Glu Ser Ser Leu Glu Ala Xaa Phe Ser Ser Ser -30 -25 -20
- Gly Ala Val Ser Gly Ala Ser Gly Phe Leu Pro Pro Ala Arg Ser Arg -15 -5 1
- The Phe Lys Ile Ile Val Ile Gly Asp Xaa Asn Val Gly Lys Thr Cys 5 10 15
- Let Thr Tyr Arg Phe Cys Ala Gly Arg Phe Pro Asp Arg Thr Glu Ala

20

25

30

Thr Ile Gly Val Asp Phe Arg Glu Arg Ala Val Glu Ile Asp Gly Glu 35 40 45

Arg Ile Lys Ile Gln Leu Trp Asp Thr Ala 50 55

- (2) INFORMATION FOR SEQ ID NO: 398:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 61 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -31..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.1 seq VSGASGFLPPARS/RI
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 398:
- Met Ala Glu Glu Met Glu Ser Ser Leu Glu Ala Ser Phe Ser Ser Ser -30 -25 -20
- Gly Ala Val Ser Gly Ala Ser Gly Phe Leu Pro Pro Ala Arg Ser Arg -15 -10 -5 1
- Ile Phe Lys Ile Ile Val Ile Gly Asp Ser Asn Val Xaa Lys Thr Cys $5 \hspace{1cm} 10 \hspace{1cm} 15$
- Leu Thr Tyr Arg Phe Cys Ala Gly Arg Phe Pro Asp Arg 20 25 30
- (2) INFORMATION FOR SEQ ID NO: 399:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 42 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE: .
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: -27..-1
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 5

seq HLSLILLKPLCLP/NN

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 399:

Met Leu Val Leu Gly Ser Pro Leu Leu Gly Pro Leu Leu Trp His Leu
-25 -20 -15

Ser Leu Ile Leu Leu Lys Pro Leu Cys Leu Pro Asn Asn Leu Pro Leu
-10 -5 1 5

Ala Leu Gly Arg Cys Leu Cys Leu His Ser 10 15

- (2) INFORMATION FOR SEQ ID NO: 400:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 82 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -55..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5 seq VLFMTTAVDLVIT/EV
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 400:

Met His Leu Leu Asp Leu Glu Ser Met Gly Lys Ser Ser Asp Gly Lys
-55 -50 -45 -40

Ser Tyr Val Ile Thr Gly Ser Trp Asn Pro Lys Ser Pro His Phe Gln
-35
-30
-25

Val Val Asn Glu Glu Thr Pro Lys Asp Lys Val Leu Phe Met Thr Thr -20 -15 -10

Ala Val Asp Leu Val Ile Thr Glu Val Gln Glu Pro Val Arg Phe Leu
-5 1 5

Leu Glu Thr Lys Val Arg Val Cys Ser Pro Asn Glu Arg Leu Phe Trp 10 20 25

Pro Ala

(2) INFORMATION FOR SEQ ID NO: 401:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 43 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8

seq VLFVFSSIPLTFL/FQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 401:

Met Glu Asn Leu Lys Asp Phe Tyr Val Leu Phe Val Phe Ser Ser Ile
-20 -15 -10

Pro Leu Thr Phe Leu Phe Gln Lys Leu Pro Phe Val Trp Ile Xaa Glu -5 1 5 10

Glu Thr Leu Glu Thr Trp Tyr Leu Lys Ser Trp
15 20

- (2) INFORMATION FOR SEQ ID NO: 402:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20 amino acids
 - (3) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -13..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8

seq LSIFSLVLPVCRM/HR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 402:

Met Pro Gln Tyr Cys Leu Ser Ile Phe Ser Leu Val Leu Pro Val Cys
-15 -10 -5

Arg Met His Arg

- (2) INFORMATION FOR SEQ ID NO: 403:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 60 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -43..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8 seq LLAFGTSCSVVLY/DP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 403:
- Met Val Ala Pro Val Leu Glu Thr Ser His Val Phe Cys Cys Pro Asn
 -40 -35 -30
- Arg Val Arg Gly Val Leu Asn Trp Ser Ser Gly Pro Arg Gly Leu Leu
 -25 -20 -15
- Ala Phe Gly Thr Ser Cys Ser Val Val Leu Tyr Asp Pro Leu Gly Cys -10 -5 1 5
- Cys Tyr Gln Leu Glu Trp Ser His Arg Pro Phe Arg
- (2) INFORMATION FOR SEQ ID NO: 404:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 71 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (3) LOCATION: -38..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8

seq LSWLITWFGHXLS/DF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 404:

Met Pro Ile Ile Asp Gln Val Asn Pro Glu Leu His Asp Phe Met Gln
-35 -30 -25

Ser Ala Glu Val Gly Thr Ile Phe Ala Leu Ser Trp Leu Ile Thr Trp
-20 -15 -10

Phe Gly His Xaa Leu Ser Asp Phe Arg His Val Val Arg Leu Tyr Asp
-5 1 5 10

Phe Phe Leu Ala Cys His Pro Leu Met Pro Ile Tyr Phe Ala Ala Val 15 20 25

Ile Val Leu Tyr Arg Glu Gln

- (2) INFORMATION FOR SEQ ID NO: 405:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 104 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -49..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.7

seq GLCVLVPCSXSXX/WR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 405:

Met Glu Thr Xaa Cys Pro Cys Cys Cys Cys Pro Cys Xaa Gly Xaa Gly -45 -40 -35

Ser Leu Xaa Xaa Lys Pro Val Tyr Glu Leu Gln Val Gln Lys Ser Val -30 -25 -20

Thr Val Gln Glu Gly Leu Cys Val Leu Val Pro Cys Ser Xaa Ser Xaa -15 -10 -5

Xaa Trp Arg Ser Trp Tyr Ser Ser Pro Pro Leu Tyr Val Tyr Trp Phe
1 5 10 15

Arg Asp Gly Glu Ile Pro Tyr Tyr Ala Glu Val·Val Ala Thr Asn Asn 20 25 30

Pro Asp Arg Arg Xaa Lys Xaa Xaa Xaa Xaa Aro Ile Pro Pro Pro 35 40 45

Trp Gly Cys Pro Glu Glu Glu Leu
50 55

- (2) INFORMATION FOR SEQ ID NO: 406:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 35 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -17..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.7

seq IYFFACFXXLTSS/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 406:

Met Ser Pro Cys Ile Tyr Phe Phe Ala Cys Phe Xaa Xaa Leu Thr Ser

Ser Ser Pro Pro His Pro Cys Pro Lys Cys Trp Pro Ser Ser Gly Ser 1 5 10 15

Ile Pro Leu

- (2) INFORMATION FOR SEQ ID NO: 407:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 33 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -27..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.7

seq VLKCLSFSXPSLP/GF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 407:

Met Gly Arg Gly Glu Arg Arg His Tyr Trp Gly Pro Lys Leu Val Leu
-25 -20 -15

Lys Cys Leu Ser Phe Ser Xaa Pro Ser Leu Pro Gly Phe Leu Trp Ser

Leu

- (2) INFORMATION FOR SEQ ID NO: 408:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 81 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -52..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.7 seq LLAKALHLLKSSC/AP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 408:

Met Ser Gln Asp Gly Gly Xaa Gly Glu Leu Lys His Met Val Met Ser
-50 -45 -40

Phe Arg Val Ser Glu Leu Gln Val Leu Leu Gly Phe Ala Gly Arg Asn -35 -30 -25

Lys Ser Gly Arg Lys His Glu Leu Leu Ala Lys Ala Leu His Leu Leu -20 - -15 -10 -5

Lys Ser Ser Cys Ala Pro Ser Val Gln Met Lys Ile Lys Glu Leu Tyr 1 5 10

Arg Arg Phe Pro Arg Lys Thr Leu Gly Pro Ser Asp Leu Ser Leu 15 20 25

Lys

- (2) INFORMATION FOR SEQ ID NO: 409:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 85 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -69..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.6

seq LGPSLSSLPSALS/LM

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 409:

Met His His Arg Met Asn Glu Met Asn Leu Ser Pro Val Gly Met Glu -65 -60 -55

Gln Leu Thr Ser Ser Ser Val Ser Asn Ala Leu Pro Val Ser Gly Ser -50 -45 -40

His Leu Gly Leu Ala Ala Ser Pro Thr His Ser Ala Ile Pro Ala Pro -35 -30 -25

Gly Leu Pro Val Ala Ile Pro Asn Leu Gly Pro Ser Leu Ser Ser Leu -20 -15 -10

Pro Ser Ala Leu Ser Leu Met Leu Pro Met Gly Xaa Gly Asp Arg Gly -5 1 5 10

Val Met Cys Gly Leu 15

- (2) INFORMATION FOR SEQ ID NO: 410:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 22 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -19..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.6 seq IWNLFSLFSTSTT/LP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 410:

Met Leu His Ser Asp Asn Ile Trp Asn Leu Phe Ser Leu Phe Ser Thr -15 -10 -5

WO 99/06550 PCT/IB98/01232

Ser Thr Thr Leu Pro Arg

- (2) INFORMATION FOR SEQ ID NO: 411:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 44 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -24..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.6 seq FHSAAGWSGGGQA/CG
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 411:

Met Gln Pro Ala Ser Pro Pro Ala Arg Trp Ser Phe His Ser Ala Ala
-20
-15
-10

Gly Trp Ser Gly Gly Gly Gln Ala Cys Gly Gly His Ser Cys Asp Gln
-5 1 5

Val Leu Ala Val Ile Glu Leu Leu Asn Pro Leu Arg 10 15 20

- (2) INFORMATION FOR SEO ID NO: 412:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 32 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.5 seq LLAGSISHMFSQA/LP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 412:

Met Cys Phe Ser Phe Leu Leu Ala Gly Ser Ile Ser His Met Phe Ser -15 -10 -5

Gln Ala Leu Pro Leu His Ser Pro Gly Leu Pro Thr Thr Asn Arg Thr $1 \hspace{1cm} 5 \hspace{1cm} 10$

- (2) INFORMATION FOR SEQ ID NO: 413:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 26 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.5 seq SILFHCSVCLFLC/QY
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 413:

Met Tyr Gly Phe Ile Ile Gly Leu Ser Ile Leu Phe His Cys Ser Val

Cys Leu Phe Leu Cys Gln Tyr His Ala Trp
-5 1 5

- (2) INFORMATION FOR SEQ ID NO: 414:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 31 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -24..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.5

seq SLLGCXLAININT/FP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 414:

Met Ser Phe Gly Xaa Ile Leu Thr Phe Arg Val Ser Leu Leu Gly Cys , -20 -15 -10

Xaa Leu Ala Ile Asn Ile Asn Thr Phe Pro Ser Asn Asn His Leu
-5 l 5

(2) INFORMATION FOR SEQ ID NO: 415:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 86 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -22..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.4 seq LGRLCAGSSGVXG/AR
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 415:
- Met Ala Val Tyr Val Gly Met Leu Arg Leu Gly Arg Leu Cys Ala Gly
 -20 -15 -10
- Ser Ser Gly Val Xaa Gly Ala Arg Ala Xaa Leu Ser Arg Ser Trp Gln
 -5 5 10
- Glu Ala Arg Leu Gln Gly Val Arg Phe Leu Ser Ser Arg Glu Val Asp 15 20 25
- Arg Met Val Ser Thr Pro Ile Gly Gly Leu Ser Tyr Val Gln Gly Cys 30 35 40
- Thr Lys Lys His Leu Asn Ser Lys Thr Val Gly Gln Cys Leu Glu Thr
 45 50 55

Thr Ala Gln Arg Val Pro 60

- (2) INFORMATION FOR SEQ ID NO: 416:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 30 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.4

seq LVSIFFFWEVTNA/FL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 416:

Met Phe Asn Thr Ile Tyr Leu Val Ile Ser Leu Val Ser Ile Phe Phe -20 -15 -10

Phe Trp Glu Val Thr Asn Ala Phe Leu Lys Ala Arg Arg Trp -5 1 5

- (2) INFORMATION FOR SEQ ID NO: 417:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 34 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -22..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.4 seq SLPLTTGSSWSLS/SQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 417:

Met Ala Leu Pro Pro Lys Gly Cys Gly Ser Leu Pro Leu Thr Thr Gly
-20 -15 -10

Ser Ser Trp Ser Leu Ser Ser Gln Ile Gly Ser Pro Ala Ile Ser Asn
-5 1 5 10

Pro Arg

- (2) INFORMATION FOR SEQ ID NO: 418:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 28 amino acids
 - (B) TYPE: AMINO ACID

- . (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -16..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.3

seq FLSWASFLAPLLR/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 418:

Met Phe Val Phe Leu Ser Trp Ala Ser Phe Leu Ala Pro Leu Leu Arg
-15 -10 -5

Ser Pro Phe Leu His Cys Leu Met Gly Met Pro Gly
1 5 10

- (2) INFORMATION FOR SEQ ID NO: 419:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 51 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -28..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.3

seq LLSCSPLXPLGKS/GF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 419:

Met Xaa Met Lys Ser Ala Asn Lys Ile Thr Leu Leu Xaa His His Leu
-25 -20 -15

Leu Ser Cys Ser Pro Leu Xaa Pro Leu Gly Lys Ser Gly Phe Ser Ser

Cys Gln Arg Leu Gly Lys Arg Ala Leu Val Phe Pro Ile Xaa Lys Xaa 5 10 15 20

Ile Ile Thr

- (2) INFORMATION FOR SEQ ID NO: 420:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 34 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -32..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.2

seq SFLLLFIVIPQTP/RP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 420:

Met Cys Asn Tyr Asn Ile Tyr Val Leu Tyr Asn Ile Gly Tyr Leu Tyr
-30 -25 -20

His Pro Lys Ser Phe Leu Leu Phe Ile Val Ile Pro Gln Thr Pro
-15 -5

Arg Pro

- (2) INFORMATION FOR SEQ ID NO: 421:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 100 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -27..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.2

seq PLLAAPLLRSLLP/RX

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 421:

Met Ala Val Ala Met Val Lys Leu Cys Glu Arg Ala Gly Leu Pro Leu
-25 -20 -15

Leu Ala Ala Pro Leu Leu Arg Ser Leu Leu Pro Arg Xaa Pro Gln Pro
-10 -5 1 5

Gly Pro Ala Gln Pro Arg Ser Val Gln Gly Gln Arg Cys Pro Ala Arg
10 15 20

His Pro Pro Gly Asn Leu Val Cys Glu Arg Gly Ala Xaa Val Asn Gly
25 30 35

Val Thr Ala Gly Ala Xaa Gly Xaa Leu Arg Gly Leu His Arg Gly Xaa 40 45 50

Arg Ala Leu Gly Cys Ser Ala His Arg Pro Xaa His Ser Ala Arg Val 55 60 65

Arg Pro Pro Ala

(2) INFORMATION FOR SEQ ID NO: 422:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 127 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -122..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.2 seq DVLLGLLKDVLLA/RP
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 422:

Met Leu Asn Val Val Arg Ala Leu Arg Xaa Pro Gln Trp Cys Ala Glu -120 -115 -110

Tyr Cys Leu Ser Ile His Tyr Gln His Gly Gly Val Ile Cys Thr Gln
-105 -100 -95

Val His Lys Gln Thr Val Val Gln Leu Ala Leu Arg Val Ala Asp Glu -90 -85 -80 -75

Met Asp Val Asn Ile Gly His Glu Val Gly Tyr Val Ile Pro Phe Glu -70 -65 -60

Asn Cys Cys Thr Asn Glu Thr Ile Leu Arg Tyr Cys Thr Asp Asp Met -55 -50 -45

Leu Gln Arg Glu Met Met Ser Asn Pro Phe Leu Gly Ser Tyr Gly Val $-40 \hspace{1.5cm} -35 \hspace{1.5cm} -30$

Ile Ile Leu Asp Asp Ile His Glu Arg Ser Ile Ala Thr Asp Val Leu
-25 -20 -15

- (2) INFORMATION FOR SEQ ID NO: 423:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 34 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -27..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.2 seq AGLCIGSTSYVHG/DI
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 423:

Met His Ala Gly Leu Glu Arg Xaa Ser Xaa Gln Lys Ala Leu Ala Gly
-25 -20 -15

Leu Cys Ile Gly Ser Thr Ser Tyr Val His Gly Asp Ile Leu Arg Thr -10 -5 1 5

Glu Arg

- (2) INFORMATION FOR SEQ ID NO: 424:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 45 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -35..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.2

seq LLGSLSLWRWSAM/EP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 424:

Met Leu Asn Gly Pro Phe Gln His Arg Asn Ser Arg Ile Met Thr His -35 -25 -25

Arg Ser Ala Glu Lys Thr Leu Leu Gly Ser Leu Ser Leu Trp Arg Trp
-15 -10 -5

Ser Ala Met Glu Pro Thr Asp Arg Cys Thr Arg Val Gly
1 5 10

- (2) INFORMATION FOR SEQ ID NO: 425:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 122 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -44..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.1 seq IAVGLTCQHVSHA/IS
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 425:

Met Arg Val Lys Asp Pro Thr Lys Ala Leu Pro Glu Lys Ala Lys Arg
-40 -35 -30

Ser Lys Arg Pro Thr Val Pro His Asp Glu Asp Ser Ser Asp Asp Ile
- -25 - -20 -15

Ala Val Gly Leu Thr Cys Gln His Val Ser His Ala Ile Ser Val Asn -10 -5 1

His Val Lys Arg Ala Ile Ala Glu Asn Leu Trp Ser Val Cys Ser Glu 5 10 15 20

Cys Leu Lys Glu Arg Arg Phe Tyr Asp Gly Gln Leu Val Leu Thr Ser 25 30 35

Asp Ile Trp Leu Cys Leu Lys Cys Gly Phe Gln Gly Cys Gly Lys Asn 40 45 50

Ser Glu Ser Gln His Ser Leu Lys His Phe Lys Ser Ser Arg Thr Glu 55 60 65

Pro His Cys Ile Ile Ile Asn Leu Ser Thr 70 75

- (2) INFORMATION FOR SEQ ID NO: 426:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 32 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -28..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4

seq FSLLALSMLKGTG/KV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 426:

Met Pro Gln Lys Gly Leu Gly Leu Gly Ile Leu Ser Gly Asp Phe
-25
-20
-15

Ser Leu Leu Ala Leu Ser Met Leu Lys Gly Thr Gly Lys Val Gly Gly -10 -5 1

- (2) INFORMATION FOR SEQ ID NO: 427:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 86 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -55..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4

seq AALCGISLSQLFP/EP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 427:

Met Ala Met Trp Asn Arg Pro Xaa Xaa Xaa Leu Pro Gln Gln Pro Leu
-55 -50 -45 -45

Xaa Ala Glu Pro Thr Ala Glu Gly Glu Pro His Leu Pro Thr Gly Arg
-35 -30 -25

Xaa Xaa Thr Glu Ala Asn Arg Phe Ala Tyr Ala Ala Leu Cys Gly Ile -20 -15 -10

Ser Leu Ser Gln Leu Phe Pro Glu Pro Glu His Ser Ser Phe Cys Thr
-5

Glu Phe Met Ala Gly Leu Val Xaa Trp Leu Glu Leu Ser Glu Ala Val 10 15 20 25

Leu Pro Thr Met Thr Ala 30

- (2) INFORMATION FOR SEQ ID NO: 428:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -19..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4

seq LLLSPWVTVPVWS/SS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 428:

Met Leu Cys Phe Gly Asp Leu Leu Ser Pro Trp Val Thr Val Pro

Val Trp Ser Ser Ser Pro Trp

- (2) INFORMATION FOR SEQ ID NO: 429:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 48 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig peptide

- (B) LOCATION: -27..-1
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 4 seq LIYFLGLAADTYF/RS
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 429:
- Met Gln Glu Asn Ala His Asn Leu Arg Leu Phe Lys Cys Leu Leu Ile
 -25 -20 -15
- Tyr Phe Leu Gly Leu Ala Ala Asp Thr Tyr Phe Arg Ser Lys Arg Lys
 -10 -5 1 5
- Pro Val Ser Phe Val Val Thr Val Xaa Xaa Gly Xaa Tyr Ala Thr Gly
 10 15 20
- (2) INFORMATION FOR SEQ ID NO: 430:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 63 amino acids
 - (3) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -59..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4

seq SVATALFPPLCIS/TG

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 430:
- Met His Thr Cys Ser Leu Pro Cys Leu Leu Phe Ala Gln Leu Leu Glu
 -55 -50 -45
- Phe Cys Ser Phe Pro Pro Asp Val Pro His Asn Cys Ala Pro Ile Val -40 -35 -30
- Ser Val Arg Pro Pro Asn Ile Val Ala Ala Phe Glu Gly Cys Ser Val
- Ala Thr Ala Leu Phe Pro Pro Leu Cys Ile Ser Thr Gly Asn Glu -10 -5 1
- (2) INFORMATION FOR SEQ ID NO: 431:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 96 amino acids
 - (B) TYPE: AMINO ACID

- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -28..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4

seq PLLGVLFFQGVYI/VF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 431:

Met Gln Gln Arg Gly Ala Ala Gly Ser Arg Gly Cys Ala Leu Phe Pro
-25 -20 -15

Leu Leu Gly Val Leu Phe Phe Gln Gly Val Tyr Ile Val Phe Ser Leu
-10 -5 1

Glu Ile Arg Ala Asp Ala His Val Arg Gly Tyr Val Gly Glu Lys Ile 5 10 15 20

Lys Leu Lys Cys Thr Phe Lys Ser Thr Ser Asp Val Thr Asp Lys Leu 25 30 35

Thr Ile Asp Trp Thr Tyr Arg Pro Pro Ser Ser Ser His Thr Val Ser 40 45 50

Ile Xaa His Tyr Gln Ser Phe Gln Tyr Pro Thr Thr Ala Gly Thr Phe
55 60 65

- (2) INFORMATION FOR SEQ ID NO: 432:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 107 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -39..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.9 seq LILNRSLPTASSS/SS
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 432:

Met Xaa Xaa Ser Ile Phe Ile Ser Glu Lys Tyr Gly Leu Cys Pro Ser

Lys Thr Pro Ile Met Lys Met Leu Pro Ser Leu Ile Leu Asn Arg Ser
-20 -15 -10

Leu Pro Thr Ala Ser Ser Ser Ser Ser Arg Lys Asp Phe Arg Leu Pro
-5 1 5

Gln Thr Arg Arg Arg Ile Ile Met Val Pro Arg Lys Glu Asp Gln Thr 10 25

Pro Leu Asn Pro Ala Ser Gln Pro Gln Ala Pro Pro Lys Pro Ile Pro 30 35 40

Ser Xaa Lys Ser Leu Glu Ala Xaa Asp Xaa Xaa Xaa Ser Gln Arg Thr 45 50 55

Xaa Arg Pro Gly Leu Ser Arg Gly Arg Ser Cys 60 65

(2) INFORMATION FOR SEQ ID NO: 433:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 75 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -20..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.9 seq FFWVVLFSAGCKV/IT
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 433:

Met Ala Phe Asp Val Ser Cys Phe Phe Trp Val Val Leu Phe Ser Ala
-20 -15 -10

Gly Cys Lys Val Ile Thr Ser Trp Asp Gln Met Tyr Ile Glu Lys Glu $1 ag{5}$

Ala Asn Lys Thr Tyr Asn Cys Glu Asn Leu Gly Leu Ser Glu Ile Pro
15 20 25

Asp Thr Leu Pro Asn Thr Thr Glu Phe Leu Glu Phe Ser Phe Asn Phe 30 35 40

Leu Pro Thr Ile His Asn Arg Thr Ser Ser Arg 45 50 55

- (2) INFORMATION FOR SEQ ID NO: 434:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 105 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -96..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.9

seq IMNLTVMLDTAXG/KX

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 434:
- Met Glu Val Ala Ala Asn Cys Ser Leu Arg Val Lys Arg Pro Leu Leu
 -95 -85
- Asp Pro Arg Phe Glu Gly Tyr Lys Xaa Ser Leu Glu Pro Leu Pro Cys
 -80 -75 -70 -65
- Tyr Gln Leu Glu Leu Asp Ala Ala Val Ala Xaa Val Lys Leu Arg Asp
 -60 -55 -50
- Asp Gln Tyr Thr Leu Glu His Met His Ala Phe Gly Met Tyr Asn Tyr
 -45 -40 -35
- Leu His Cys Asp Ser Trp Tyr Gln Asp Ser Val Tyr Tyr Ile Asp Thr -30 -25 -20
- Leu Gly Arg Ile Met Asn Leu Thr Val Met Leu Asp Thr Ala Xaa Gly -15 -10 -5
- Lys Xaa Arg Glu Val Phe Arg Leu Leu
- (2) INFORMATION FOR SEQ ID NO: 435:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 95 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -39..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.9 seq VLAIGLLHIVLLS/IP
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 435:

Met Asn Val Gly Thr Ala His Xaa Xaa Val Asn Pro Asn Thr Arg Val -35 -30 -25

Met Asn Ser Arg Gly Ile Trp Leu Ser Tyr Val Leu Ala Ile Gly Leu -20 -15 -10

Leu His Ile Val Leu Leu Ser Ile Pro Phe Val Ser Val Pro Val Val -5 1 5

Trp Thr Leu Thr Asn Leu Ile His Asn Met Gly Met Tyr Ile Phe Leu 10 20 25

His Thr Val Lys Gly Xaa Pro Phe Glu Thr Pro Asp Gln Gly Lys Ala 30 35 40

Arg Leu Leu Xaa His Xaa Xaa Ala Asp Gly Leu Trp Gly Pro Val

- (2) INFORMATION FOR SEQ ID NO: 436:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 48 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.9 seq SWWTLLSSSPSFM/IS
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 436:

Met Glu Asn Phe Asn Met Tyr Lys Asn Lys Ser Trp Trp Thr Leu Leu -20 -15 -10

Ser Ser Ser Pro Ser Phe Met Ile Ser Phe Val Ser Ser Val Leu Pro

Val Leu Leu Thr Ile Ser Arg Phe Ile Leu Lys Gln Ile Pro Asp Gln 10 20 25

- (2) INFORMATION FOR SEQ ID NO: 437:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 70 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -39..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.9

seq VLAIGLLHIVLLS/IP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 437:

Met Asn Val Gly Thr Xaa His Ser Glu Val Asn Pro Asn Thr Arg Val
-35
-30
-25

Met Asn Ser Arg Gly Ile Trp Leu Ser Tyr Val Leu Ala Ile Gly Leu
-20 -15 -10

Leu His Ile Val Leu Leu Ser Ile Pro Phe Val Ser Val Pro Val Val

Trp Thr Leu Thr Asn Leu Ile His Asn Met Gly Met Tyr Ile Phe Leu 10 20 25

Tyr Thr Val Lys Gly Thr

- (2) INFORMATION FOR SEQ ID NO: 438:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 49 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -14..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.8

WO 99/06550 PCT/IB98/01232

seq AAASAVSVLLVAA/ER

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 438:

Met Ala Ala Ser Ala Val Ser Val Leu Leu Val Ala Ala Glu Arg

Asn Arg Trp His Arg Leu Pro Ser Leu Leu Leu Pro Pro Arg Thr Trp 5 10 15

Val Trp Arg Gln Arg Thr Met Lys Tyr Thr Thr Ala Thr Gly Arg Asn 20 25 30

Met 35

- (2) INFORMATION FOR SEQ ID NO: 439:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 95 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -44..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.8 seq SGSGLSWARLSQS/RS
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 439:
- Met Ala Tyr Ser Lys Ala Ser Gly Ser Pro Val Leu Ser Gln Ala Val -40 -35 -30
- Pro Gly Glu Asn Ala Ser His Arg Arg Gly Ser Ala Asp Leu Gly Ser -25 -20 -15
- Gly Ser Gly Leu Ser Trp Ala Arg Leu Ser Gln Ser Arg Ser Glu Ile -10 -5 1
- His Ser Ala Gly Pro Pro His Leu Gly Gly Arg Thr Asn Gly Pro Glu
 5 10 15 20
- Phe Pro Ala Leu Ser Tyr Ser Ser Gln Leu Leu Ser Leu Ala Gln Leu 25 30 35
- Arg Gly Arg Gly Ile Thr Glu Val Ser Glu Lys Ser Pro Leu Ile 40 45 50

- (2) INFORMATION FOR SEQ ID NO: 440:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 95 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -37..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.8

seq RPVLLHLHQTAHA/DE

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 440:
- Met Lys Pro Arg Arg Asn Leu Glu Glu Asp Asp Tyr Leu His Lys Asp
 -35
 -30
 -25
- Thr Gly Glu Thr Ser Met Leu Lys Arg Pro Val Leu Leu His Leu His -20 -15 -10
- Gln Thr Ala His Ala Asp Glu Phe Asp Cys Pro Ser Glu Leu Gln His
 -5 1 5 10
- Thr Gln Glu Leu Phe Pro Gln Trp His Leu Pro Ile Lys Ile Ala Ala 15 20 25
- Ile Ile Ala Ser Leu Thr Phe Leu Tyr Thr Leu Leu Arg Glu Val Ile 30 35 40
- His Pro Leu Ala Thr Ser His Gln Gln Tyr Phe Tyr Lys Ile Gln 45 50 55
- (2) INFORMATION FOR SEQ ID NO: 441:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 39 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -19..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 3.7

seq IPCAHMLVCPTIG/DI

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 441:

Met Ile Ile Cys Tyr Asp Ile Pro Cys Ala His Met Leu Val Cys Pro
-15 -10 -5

Thr Ile Gly Asp Ile Lys Phe Asp His Leu Met Lys Trp Tyr Pro Ser $1 \hspace{1cm} 5 \hspace{1cm} 10$

Asp Phe Ser Thr Glu Arg Leu 15 20

- (2) INFORMATION FOR SEQ ID NO: 442:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 70 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -19..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.7

seq STLASVPPAATFG/AD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 442:

Met Tyr Ser Ser Glu Asp Ser Thr Leu Ala Ser Val Pro Pro Ala Ala
-15 -10 -5

Thr Phe Gly Ala Asp Asp Leu Val Leu Thr Leu Ser Asn Pro Gln Met $1 \hspace{1cm} 5 \hspace{1cm} 10$

Ser Leu Glu Gly Thr Glu Lys Ala Ser Trp Leu Gly Glu Gln Pro Gln
15

Phe Trp Ser Lys Thr Gln Val Leu Asp Trp Ile Ser Tyr Gln Val Glu 30 40 45

Lys Asn Lys Tyr Asp Ala

- (2) INFORMATION FOR SEQ ID NO: 443:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 70 amino acids

- (B) TYPE: AMINO ACID (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -65..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.7

seq QLEGLNWLRFSWA/QG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 443:

Met Gly Glu Asp Pro Xaa Gln Pro Arg Lys Tyr Lys Lys Xaa Lys Xaa

Glu Leu Gln Gly Asp Xaa Pro Pro Ser Ser Pro Thr Asn Asp Pro Thr -45 -40

Val Lys Tyr Glu Thr Gln Pro Arg Phe Ile Thr Ala Thr Gly Gly Thr -25

Leu His Met Tyr Gln Leu Glu Gly Leu Asn Trp Leu Arg Phe Ser Trp -10

Ala Gln Gly Thr Xaa Gly

- (2) INFORMATION FOR SEQ ID NO: 444:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 75 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -42..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.7 seq LLGCLQCCWLQSG/RA
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 444:

Met Phe Tyr Val Ala Met Thr Lys Thr His Lys Arg Ile Arg Ser Leu -40 -35

Cys Asn Ile His His Gly Leu Phe Gln Phe Thr Gln Gln Leu Leu Gly -25 -15

Cys Leu Gln Cys Cys Trp Leu Gln Ser Gly Arg Ala Pro Ala Thr Tyr -10 -5 1 5

Tyr Leu Val Glu Ser Ile Glu Lys Ser Ala His Gly Ser Val Leu Xaa

Thr Tyr Asp Gln Thr Gln Thr Arg Ile Gly Arg

- (2) INFORMATION FOR SEQ ID NO: 445:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 62 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -60..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.7 seq XTCASXNPSQCLA/AF
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 445:

Met Val Ser Pro Lys Asp Leu Pro Leu Val Leu Gln Asp Ile Lys
-60 -55 -50 -45

Val Pro Ser Ser Met Thr Gly Ser His Ala Gly Asn Pro His Ile Glu
-40 -35 -30

Arg Asn Asp Leu Pro Arg His Gly Ser Pro Gln Phe Phe Thr Gly Xaa -25 -20 -15

Thr Cys Ala Ser Xaa Asn Pro Ser Gln Cys Leu Ala Ala Phe
-10 -5 1

- (2) INFORMATION FOR SEQ ID NO: 446:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 26 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.6

seq FXSLFCLYFSCFL/HI

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 446:

Met Glu Phe Xaa Ser Leu Phe Cys Leu Tyr Phe Ser Cys Phe Leu His -15 -10 -5 1

Ile Ile Tyr Phe Xaa Ser Cys Phe Leu Tyr
5

- (2) INFORMATION FOR SEQ ID NO: 447:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 66 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -45..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.6

seq ALLELIDSPECLS/KC

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 447:

Met Ala Leu His Phe Gln Ser Leu Ala Glu Leu Glu Xaa Leu Cys Thr -45 -40 -35 -30

His Leu Tyr Ile Gly Thr Asp Leu Thr Gln Arg Ile Glu Ala Glu Lys
-25
-20
-15

Ala Leu Leu Glu Leu Ile Asp Ser Pro Glu Cys Leu Ser Lys Cys Gln -10 -5 1

Leu Leu Glu Gln Gly Thr Thr Ser Tyr Ala Gln Leu Leu Ala Ala 5 10 15

Thr Xaa

20

(2) INFORMATION FOR SEQ ID NO: 448:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 40 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -27..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.6

seq LTLLLITPSPSPL/LF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 448:

Met Arg Thr Leu Phe Gly Ala Val Arg Ala Pro Phe Ser Ser Leu Thr
-25 -20 -15

Leu Leu Leu Ile Thr Pro Ser Pro Ser Pro Leu Leu Phe Asp Arg Gly
-10 -5 1 5

Leu Ser Leu Arg Ser Ala Met Ser 10

- (2) INFORMATION FOR SEQ ID NO: 449:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 44 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -41..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.6

seq AVSSLIAVGTSHG/LA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 449:

Met Arg His Ser Leu Leu Lys Gly Ile Ser Ala Gln Ile Val Ser Ala
-40 -35 -30

Ala Asp Lys Val Asp Ala Gly Leu Pro Thr Ala Ile Ala Val Ser Ser
-25 -15 -10

Leu Ile Ala Val Gly Thr Ser His Gly Leu Ala Gly
•-5

- (2) INFORMATION FOR SEQ ID NO: 450:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.5 seq LSCFIFFYISSLC/CF
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 450:

Met Thr Leu Ser Cys Phe Ile Phe Phe Tyr Ile Ser Ser Leu Cys Cys -15 -5 1

Phe Leu Ser Tyr Pro Thr Arg

- (2) INFORMATION FOR SEQ ID NO: 451:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 47 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.5

seq LCFLLPHHRLQEA/RQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 451:

Met Ile Leu Cys Phe Leu Leu Pro His His Arg Leu Gln Glu Ala Arg
-15 -5 1

Gln Ile Gln Val Leu Lys Met Leu Pro Arg Glu Lys Leu Arg Arg 10 15

Arg Arg Glu Lys Thr Asn Lys Trp Glu Lys Arg Lys Gly Ser Gly 25 30

- (2) INFORMATION FOR SEQ ID NO: 452:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 24 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -14..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.5 seq FSLFALNMPLGFC/VY

.

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 452:

Met Phe Ser Leu Phe Ala Leu Asn Met Pro Leu Gly Phe Cys Val Tyr

Val Ile Phe Lys Ile His Asp Trp 5 10

- (2) INFORMATION FOR SEQ ID NO: 453:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 47 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -31..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.5

seq SVWGVLPPPACSA/DL

. . .

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 453:

Met Ala Ser Ser Pro Gly Val Ala Met His Ser Leu Trp Ala Thr Ile

His Thr Ser Val Trp Gly Val Leu Pro Pro Pro Ala Cys Ser Ala Asp
-15 -5 1

Leu Leu Phe Ser Asn Ala Cys Leu Leu Pro His Glu Ile His Leu
5 10 15

- (2) INFORMATION FOR SEQ ID NO: 454:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 86 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -45..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.5 seq LPRLLSLSQHSES/WI
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 454:

Met Ser Gln Glu Gly Ala Val Pro Ala Ser Ala Val Pro Leu Glu Glu -45 -35 -36

Leu Ser Ser Trp Pro Glu Glu Leu Cys Arg Arg Glu Leu Pro Ser Val

Leu Pro Arg Leu Leu Ser Leu Ser Gln His Ser Glu Ser Trp Ile Glu
-10 -5 1

His Ile Gln Ile Leu Lys Ile Ile Val Glu Met Phe Leu Pro His Met $5 \hspace{1cm} 10 \hspace{1cm} 15$

Asn His Leu Thr Leu Glu Gln Thr Phe Phe Ser Gln Val Leu Pro Lys 20 25 30 35

Thr Val Lys Leu Phe Asp

4 (

(2) INFORMATION FOR SEQ ID NO: 455:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 39 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -36..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.5

seq AAVVFAVVLSIHA/TV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 455:

Met Thr Arg Glu Cys Pro Ser Pro Ala Pro Gly Pro Gly Ala Pro Leu
-35 -30 -25

Ser Gly Ser Val Leu Ala Glu Ala Ala Val Val Phe Ala Val Val Leu
-20 -15 -10 -5

Ser Ile His Ala Thr Val Trp

- (2) INFORMATION FOR SEQ ID NO: 456:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 85 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 14.8

seq LLWWALLLGLAQA/CP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 456:

Met Gln Glu Leu His Leu Leu Trp Trp Ala Leu Leu Leu Gly Leu Ala -15 -10 -5

Gln Ala Cys Pro Glu Pro Cys Asp Cys Gly Glu Lys Tyr Gly Phe Gln
1 5 10

Ile Ala Asp Cys Ala Tyr Arg Asp Leu Glu Ser Val Pro Pro Gly Phe
15 20 25 30

Pro Ala Asn Val Thr Thr Leu Ser Leu Ser Ala Asn Arg Leu Pro Gly
•35 40 45

Leu Pro Glu Gly Ala Phe Arg Glu Val Pro Leu Leu Gln Ser Leu Trp 50 55 60

Leu Ala His Asn Glu 65

- (2) INFORMATION FOR SEQ ID NO: 457:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 106 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 13.6 seq LLLLALCATGAQG/LY
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 457:
- Met Gly Arg Gln Ala Leu Leu Leu Leu Ala Leu Cys Ala Thr Gly Ala
 -15 -10 -5
- Gin Gly Leu Tyr Phe His Ile Gly Glu Thr Glu Lys Arg Cys Phe Ile
 1 5 10
- Glu Glu Ile Pro Asp Glu Thr Met Val Ile Gly Asn Tyr Arg Thr Gln
 15 20 25 30
- Met Trp Asp Lys Gln Lys Glu Val Phe Leu Pro Ser Thr Pro Gly Leu 35 40 45
- Gly Met His Val Glu Val Lys Asp Pro Asp Gly Lys Val Val Leu Ser 50 55 60
- Arg Gln Tyr Gly Ser Glu Gly Arg Phe Thr Phe Thr Ser His Xaa Xaa 65 70 75
- Gly Asp His Gln Ile Cys Leu His Cys Gly
- (2) INFORMATION FOR SEQ ID NO: 458:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 40 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 12.7

seq ILFLLSWSGPLQG/QQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 458:

Met Gly Pro Ser Thr Pro Leu Leu Ile Leu Phe Leu Leu Ser Trp Ser
-20 -15 -10

Gly Pro Leu Gln Gly Gln Gln His His Leu Val Glu Tyr Met Glu Arg -5 10

Arg Leu Ala Ala Leu Glu Glu Arg

- (2) INFORMATION FOR SEQ ID NO: 459:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 78 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -27..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.8

seq LLLLCPLSRGCCP/LL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 459:

Met Ser Cys Arg Glu Leu Thr His Arg Pro Cys Ser Pro His Leu Leu
-25 -20 -15

Leu Leu Cys Pro Leu Ser Arg Gly Cys Cys Pro Leu Leu Ser Xaa -10 -5 !

Pro Leu Xaa Gly Val Asn Leu Glu Ser Ile Leu Ser Leu Thr Leu Pro 10 15 20

Pro Ser Pro Ser Ser Val Gly Leu Ser Pro Ser Val Thr Xaa Leu Thr
25 30 35

Thr Ser Pro Val Ser Leu His Phe Ala Ser Xaa Leu Ala Gly
40 45 50

(2) INFORMATION FOR SEQ ID NO: 460:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 121 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -22..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.5

seq AALLLGLMMVVTG/DE

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 460:
- Met Gly Trp Thr Met Arg Leu Val Thr Ala Ala Leu Leu Gly Leu -20 -15 -10
- Met Met Val Val Thr Gly Asp Glu Asp Glu Asp Ser Pro Cys Ala His
 -5 1 5 10
- Glu Ala Leu Leu Asp Glu Asp Thr Leu Phe Cys Gln Gly Leu Glu Val 15 20 25
- Phe Tyr Pro Glu Leu Gly Asn Ile Gly Cys Lys Val Val Pro Asp Cys $30 \hspace{1cm} 35 \hspace{1cm} 40$
- Xaa Asn Tyr Arg Gln Lys Ile Thr Ser Trp Met Glu Pro Ile Val Lys
 45 50 55
- Phe Pro Gly Ala Val Asp Gly Ala Thr Tyr Ile Leu Val Met Val Asp 60 65 70
- Pro Asp Ala Pro Ser Arg Ala Glu Pro Arg Gln Arg Phe Trp Arg His 75 80 85 90
- Trp Leu Val Thr Asp Ile Lys Gly Ala

(2) INFORMATION FOR SEQ ID NO: 461:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 33 amino acids

(B) TYPE: AMINO ACID

(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: PROTEIN

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: -24..-1

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 7.3

seq VHLLSLCSGKVYA/RM

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 461:

Met Lys Phe Leu Ile Phe Ala Phe Phe Gly Gly Val His Leu Leu Ser

Leu Cys Ser Gly Lys Val Tyr Ala Arg Met Ala Ser Leu Arg Gly Leu -5 1 5

Gly

- (2) INFORMATION FOR SEQ ID NO: 462:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 89 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -29..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.1

seq LIFLCGAALLAVG/IW

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 462:

Met Gln Cys Phe Ser Phe Ile Lys Thr Met Met Ile Leu Phe Asn Leu -25 -20 -15

Leu Ile Phe Leu Cys Gly Ala Ala Leu Leu Ala Val Gly Ile Trp Val $^{-10}$ $^{-5}$ 1

Ser Ile Asp Gly Ala Ser Phe Leu Lys Ile Phe Gly Pro Leu Ser Ser 5 10 15

Ser Ala Met Gln Phe Val Asn Val Gly Tyr Phe Leu Ile Ala Ala Gly 20 25 30 35

Glu Ser Lys Cys Ala Leu Val Thr Phe 55 60

- (2) INFORMATION FOR SEQ ID NO: 463:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 51 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -28..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.6 seq IVSLLGFVATVTL/IP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 463:

Met Trp Ala Phe Ser Glu Leu Pro Met Pro Leu Leu Ile Asn Leu Ile
-25
-20
-15

Val Ser Leu Leu Gly Phe Val Ala Thr Val Thr Leu Ile Pro Ala Phe
-10 -5

Arg Gly His Phe Ile Ala Ala Arg Leu Cys Gly Gln Asp Leu Asn Lys 5 10 15 20

Thr Ser Gln

- (2) INFORMATION FOR SEQ ID NO: 464:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 85 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: -19..-1

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 6.3

seq VLMRLVASAYSIA/QK

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 464:

Met Ala Ser Ser Asn Thr Val Leu Met Arg Leu Val Ala Ser Ala Tyr -15 -10 -5

Ser Ile Ala Gln Lys Ala Gly Met Ile Val Arg Arg Val Ile Ala Glu $1 \hspace{1cm} 5 \hspace{1cm} 10$

Gly Asp Leu Gly Ile Val Glu Xaa Thr Cys Ala Thr Asp Leu Gln Thr 15 20 25

Lys Ala Asp Arg Leu Ala Gln Met Xaa Ile Cys Ser Ser Leu Ala Arg 30 3540

Lys Phe Pro Lys Leu Thr Ile Ile Gly Glu Glu Asp Leu Pro Ser Xaa 50 55 60

Glu Val Asp Gln Glu 65

(2) INFORMATION FOR SEQ ID NO: 465:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 71 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -24..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq VHLLSLCSGKAIC/KN

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 465:

Met Lys Phe Leu Ile Phe Ala Phe Phe Gly Gly Val His Leu Leu Ser

Let Cys Ser Gly Lys Ala Ile Cys Lys Asn Gly Ile Ser Lys Arg Thr $^{-5}$ 1 5

Phe Glu Glu Ile Lys Glu Glu Ile Ala Ser Cys Gly Asp Val Ala Lys
10 20

Ala Ile Ile Asn Leu Ala Val Tyr Gly Lys Ala Gln Asn Arg Ser Tyr 25 30 35 40

Xaa Arg Leu Ala Leu Leu Val

- (2) INFORMATION FOR SEQ ID NO: 466:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 118 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -51..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq ALXVLPLLGLHEA/AS

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 466:
- Met Ala Asp Thr Thr Pro Asn Gly Pro Gln Gly Ala Gly Ala Val Gln
 -50 -45 -40

Phe Met Met Thr Asn Lys Leu Asp Thr Ala Met Trp Leu Ser Arg Leu
-35 -25 -20

Phe Thr Val Tyr Cys Ser Ala Leu Xaa Val Leu Pro Leu Leu Gly Leu
-15 -10 -5

His Glu Ala Ala Ser Phe Tyr Gln Arg Ala Leu Leu Ala Asn Ala Leu 1 5 10

Thr Ser Ala Leu Arg Leu His Gln Arg Leu Pro His Phe Gln Leu Ser 15 20 25

Arg Ala Phe Leu Ala Gln Ala Leu Leu Glu Asp Ser Cys His Tyr Leu 30 40 45

Leu Tyr Ser Leu Ile Phe Val Asn Ser Tyr Pro Val Thr Met Ser Ile 50 55 60

Phe Pro Val Leu Leu Phe

65

- (2) INFORMATION FOR SEQ ID NO: 467:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 73 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -24..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.5 seq XVLVLSVVXXAMA/AF
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 467:
- Met Arg Phe Arg His Phe Xaa Lys Xaa Ile Gly Xaa Val Leu Val Leu -20 -15 -10
- Ser Val Val Xaa Xaa Ala Met Ala Ala Phe Ala Val Xaa Pro Gln Gly -5 1 5
- Pro Ala Leu Xaa Ser Glu Pro Xaa Xaa Xaa Gly Ser Pro Thr Ser Pro 10 15 20
- Lys Pro Gly Val Asn Ala Gln Phe Leu Pro Gly Phe Leu Met Gly Xaa 25 30 35 40

Leu Pro Ala Pro Val Thr Pro Gln Pro 45

- (2) INFORMATION FOR SEQ ID NO: 468:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 161 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -40..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.2 seq LCVEFASVASCDA/AV
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 468:

PCT/IB98/01232 WO 99/06550

Met Glu Leu Gly Ser Cys Leu Glu Gly Gly Arg Glu Ala Ala Glu Glu

Glu Gly Glu Pro Glu Val Lys Lys Arg Arg Leu Leu Cys Val Glu Phe

Ala Ser Val Ala Ser Cys Asp Ala Ala Val Ala Gln Cys Phe Leu Ala

Glu Asn Asp Trp Glu Met Glu Arg Ala Leu Asn Ser Tyr Phe Glu Pro 15

Pro Val Glu Glu Ser Ala Leu Glu Arg Arg Pro Glu Thr Ile Ser Glu 35

Pro Lys Thr Tyr Val Asp Leu Thr Asn Glu Glu Thr Thr Asp Ser Thr

Thr Ser Lys Ile Ser Pro Ser Glu Asp Thr Gln Gln Glu Asn Gly Ser

Met Phe Ser Leu Ile Thr Trp Asn Ile Asp Gly Leu Asp Leu Asn Asn

Leu Ser Glu Arg Ala Arg Gly Val Cys Ser Tyr Leu Ala Leu Tyr Ser

Pro Asp Val Ile Phe Leu Gln Glu Val Ile Pro Pro Tyr Tyr Ser Tyr 115

Leu

(2) INFORMATION FOR SEQ ID NO: 469:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 144 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -122..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5 seq RLVVVSVSPQSRA/SL
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 469:

Met Ala Ser Pro Phe Ser Gly Ala Leu Gln Leu Thr Asp Leu Asp Asp -120 -115 -110

Phe Ile Gly Pro Ser Gln Glu Cys Ile Lys Pro Val Lys Val Glu Lys
-105 -100 -95

- Arg Ala Gly Ser Gly Val Ala Lys Ile Arg Ile Glu Asp Asp Gly Ser -90 -85 -80 -75
- Tyr Phe Gln Ile Asn Gln Asp Gly Xaa Thr Arg Arg Leu Glu Lys Ala -70 -65 -60
- Lys Val Ser Leu Asn Tyr Cys Xaa Ala Cys Ser Gly Cys Ile Thr Ser
 -55 -50 -45
- Ala Glu Thr Val Leu Ile Thr Gln Gln Ser His Glu Glu Leu Lys Lys -40 -35 -30
- Val Leu Asp Ala Asn Lys Met Ala Ala Pro Ser Gln Gln Arg Leu Val -25 -20 -15
- Val Val Ser Val Ser Pro Gln Ser Arg Ala Ser Leu Ala Ala Arg Phe
 -10 -5 1 5
- Gln Leu Xaa Pro Thr Asp Thr Ala Arg Lys Leu Thr Ser Phe Phe Lys 10 15 20
- (2) INFORMATION FOR SEQ ID NO: 470:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 84 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -44..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8

seq SLVAELLLGAGSG/SH

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 470:
- Met Gly Pro Val Pro Thr Ala Val Ala Gly Ala Gly Ser Arg Leu Val -40 -35 -30
- Lys Pro Ser Gln Thr Leu Ser Leu Thr Cys Ala Val Ser Gly Gly Ser -25 -20 -15
- Leu Val Ala Glu Leu Leu Gly Ala Gly Ser Gly Ser His Leu Gly -10 -5 l
- Arg Ala Trp Ser Gly Leu Gly Ser Ser Ile Ile Glu Ala Ile Val Gly
 5 10 15 20

PCT/IB98/01232 WO 99/06550

Val Leu Leu Thr Ile Arg Pro Ser Arg Leu Glu Pro Pro Tyr His Trp 30 25

Thr Ser Pro Ala

- (2) INFORMATION FOR SEQ ID NO: 471:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 88 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.4

seq QFILLGTTSVVTA/AL

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 471:
- Met Glu Ser Gly Gly Arg Pro Ser Leu Cys Gln Phe Ile Leu Leu Gly
- Thr Thr Ser Val Val Thr Ala Ala Leu Tyr Ser Val Tyr Arg Gln Lys
- Ala Arg Val Ser Gln Glu Leu Lys Gly Ala Lys Lys Val His Leu Gly
- Glu Asp Leu Lys Ser Ile Leu Ser Glu Xaa Pro Gly Lys Cys Val Pro
- Tyr Ala Val Ile Glu Gly Ala Val Arg Ser Val Lys Glu Thr Leu Asn

Ser Gln Phe Val Glu Asn Cys Lys 60

- (2) INFORMATION FOR SEQ ID NO: 472:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 21 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -19..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.1

seq IYIICFXLPPLFS/FN

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 472:

Met Gln Val Cys Arg Cys Ile Tyr Ile Ile Cys Phe Xaa Leu Pro Pro -15 -10 -5

Leu Phe Ser Phe Asn

- (2) INFORMATION FOR SEQ ID NO: 473:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 50 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.1

seq QRLLLRFLASVIS/RK

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 473:

Met Ala Gln Arg Leu Leu Arg Phe Leu Ala Ser Val Ile Ser Arg
-15 -5 1

Lys Pro Ser Gln Gly Gln Trp Ala Thr Pro His Phe Gln Ser Pro Ala 5 10 15

Asp Pro Thr Met Gln Ser Trp Trp Pro Asp Cys Asn Thr Gln Pro Ser 20 25 30

Pro Asp

35

(2) INFORMATION FOR SEQ ID NO: 474:

-30

-25

-20

455 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 46 amino acids (B) TYPE: AMINO ACID (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: PROTEIN (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Cancerous prostate (ix) FEATURE: (A) NAME/KEY: sig peptide (B) LOCATION: -40..-1 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.9 seg FLWLITRPQPVLP/LL (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 474: Met Leu Phe Ile Phe Asn Phe Leu Phe Ser Pro Leu Pro Thr Pro Ala -30 Leu Ile Cys Ile Leu Thr Phe Gly Ala Ala Ile Phe Leu Trp Leu Ile -20 -15 Thr Arg Pro Gln Pro Val Leu Pro Leu Leu Asp Leu Asn Xaa -5 1 (2) INFORMATION FOR SEQ ID NO: 475: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 49 amino acids (B) TYPE: AMINO ACID (D) TOPOLOGY: LINEAR (ii) MOLECULE TYPE: PROTEIN (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: Hypertrophic prostate (ix) FEATURE: (A) NAME/KEY: sig_peptide (B) LOCATION: -46..-1 (C) IDENTIFICATION METHOD: Von Heijne matrix (D) OTHER INFORMATION: score 3.9 seq SHMLQLLPSKALC/LF (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 475: Met Tyr Pro Lys Trp Glu Ala Pro Val Thr Phe Cys Gln Leu Lys Arg -45 Glu Lys Asp Pro Pro His Pro Ala His Ser Pro Phe Leu Gln Pro Ara Phe Ser His Met Leu Gln Leu Leu Pro Ser Lys Ala Leu Cys Leu Phe
-10 -5 1

Phe

- (2) INFORMATION FOR SEQ ID NO: 476:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 60 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -44..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.7 seq LAERLGLFEELWA/AQ
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 476:

Met Ala Leu Tyr Gln Arg Trp Arg Cys Leu Arg Leu Gln Gly Leu Gln
-40 -35 -30

Ala Cys Arg Leu His Thr Ala Val Val Ser Thr Pro Pro Arg Trp Leu
-25 -20 -15

Ala Glu Arg Leu Gly Leu Phe Glu Glu Leu Trp Ala Ala Gln Val Lys

Arg Leu Ala Ser Met Ala Gln Lys Glu Pro Gln Thr
5 10 15

- (2) INFORMATION FOR SEQ ID NO: 477:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 61 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 13.8 seq XGLLLFLLPGSLG/AE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 477:

Met Gly Val Pro Arg Pro Gln Pro Trp Ala Xaa Gly Leu Leu Phe
-20 -15 -10

Leu Leu Pro Gly Ser Leu Gly Ala Glu Ser His Leu Ser Leu Leu Tyr
-5 1 5

His Leu Thr Ala Val Ser Ser Pro Ala Pro Gly Thr Pro Ala Phe Trp 10 15 20 25

Val Ser Gly Trp Leu Gly Pro Gln Gln Tyr Pro Ser Xaa 30 35

- (2) INFORMATION FOR SEQ ID NO: 478:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 109 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -45..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 13.4 seq LVLALXLVSAALS/SV
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 478:

Met Ala Ala Ala Val Pro Lys Arg Met Arg Gly Pro Ala Gln Ala Lys
-45 -35 -30

Leu Leu Pro Gly Ser Ala Ile Gln Ala Leu Val Gly Leu Ala Arg Pro
-25 -20 -15

Leu Val Leu Ala Leu Xaa Leu Val Ser Ala Ala Leu Ser Ser Val Val
-10 -5 1

Ser Arg Thr Asp Ser Pro Ser Pro Thr Val Leu Asn Ser His Iie Ser 5 10 15

Thr Pro Asn Val Asn Ala Leu Thr His Glu Asn Gln Thr Lys Pro Ser 20 25 30 35

Ile Ser Gln Ile Ser Thr Thr Leu Pro Pro Xaa Xaa Ser Thr Lys Xaa 40 45 50

Ser Gly Gly Ala Xaa Val Val Pro His Pro Ser Pro Gly 55 60

- (2) INFORMATION FOR SEQ ID NO: 479:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 66 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -29..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 13

seq LLLVLLLVTRXRS/MP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 479:

Met Trp Leu Trp Glu Asp Gln Gly Gly Leu Leu Gly Pro Phe Ser Phe
-25
-15

Leu Leu Val Leu Leu Val Thr Arg Xaa Arg Ser Met Pro Ala-10 -5 1

Ser Ser Pro Ala Ala Ser Ser Phe Tyr Cys Ala Ser Ser Ala Xaa Ser $\frac{5}{10}$ 15

Arg Cys Pro Leu Ala Gly Pro Cys Arg Cys Ser Ser Pro Gly Thr Ala 20 25 30 35

Phe Leu

- (2) INFORMATION FOR SEQ ID NO: 480:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 82 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 11.6 seq LLLLVQLLRFLRA/DG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 480:

Met Asn Trp Glu Leu Leu Leu Trp Leu Leu Val Leu Cys Ala Leu Leu -25 -20 -15

Leu Leu Val Gln Leu Leu Arg Phe Leu Arg Ala Asp Gly Asp Leu -10 -5 1

Thr Leu Leu Trp Ala Glu Trp Gln Gly Arg Arg Pro Glu Trp Glu Leu 5 10 15 20

Thr Asp Met Val Val Trp Val Thr Gly Ala Ser Ser Gly Ile Gly Glu 25 30 35

Glu Leu Ala Tyr Gln Leu Ser Lys Leu Gly Val Ser Leu Val Leu Ser 40 45 50

Ala Arg

- (2) INFORMATION FOR SEQ ID NO: 481:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 56 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -20..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 11.2

seq AFLLLVALSYTLA/RD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 481:

Met Glu Lys Ile Pro Val Ser Ala Phe Leu Leu Leu Val Ala Leu Ser -20 -15 -10 -5

Tyr Thr Leu Ala Arg Asp Thr Thr Val Lys Pro Gly Ala Lys Lys Asp $1 \hspace{1cm} 5$

Thr Lys Asp Ser Arg Pro Lys Leu Pro Gln Thr Leu Ser Arg Gly Trp
15 20 25

Gly Asp Gln Leu Ile Trp Thr Arg

- (2) INFORMATION FOR SEQ ID NO: 482:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) EENGTH: 62 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -40..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 11.2 seq AFLLLVALSYTLA/RD
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 482:

Met Ser Asn Tyr Thr Asp Ala Glu Ser Ser Phe Ser Lys Gln Glu Ile
-40 -35 -30 -25

Ile Arg Val Ala Met Glu Lys Ile Pro Val Ser Ala Phe Leu Leu -20 -15 -10

Val Ala Leu Ser Tyr Thr Leu Ala Arg Asp Thr Thr Val Lys Pro Gly
-5 1 5

Ala Lys Lys Asp Thr Lys Asp Ser Arg Pro Lys Pro Pro Arg
10 15 20

- (2) INFORMATION FOR SEQ ID NO: 483:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 108 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -53..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.6

seq FILLLIFIAEVAA/AV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 483:

Met Gln Phe Xaa Thr Trp Ala Thr Ser Ser Ser Gln Pro Ala Leu Trp
-50 -45 -40

Ser Leu Leu Val Ser Trp Ala Ala Met Val Leu Arg Leu Arg Ser -35 -30 -25

Lys Cys Ala Leu Val Thr Phe Phe Phe Ile Leu Leu Leu Ile Phe Ile
-20 -15 -10

Ala Glu Val Ala Ala Ala Val Val Ala Leu Val Tyr Xaa Thr Met Xaa -5 1 10

Glu His Phe Leu Thr Leu Leu Val Val Pro Ala Ile Lys Lys Asp Tyr 15 20 25

Gly Ser Gln Glu Asp Phe Thr Gln Val Xaa Asn Thr Thr Met Lys Gly $30 \hspace{1cm} 35 \hspace{1cm} 40$

Leu Lys Cys Cys Gly Phe Thr Asn Tyr Thr Asp Trp
45 50 55

(2) INFORMATION FOR SEQ ID NO: 484:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 111 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -28..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.5

seq LLLLVHLLRFLRA/DG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 484:

Met Asn Trp Glu Leu Leu Leu Trp Leu Leu Val Leu Cys Ala Leu Leu -25 -20 -15

Leu Leu Val His Leu Leu Arg Phe Leu Arg Ala Asp Gly Asp Leu
-10 -5 1

Thr Leu Leu Trp Ala Glu Trp Gln Gly Arg Arg Pro Glu Trp Glu Leu 5 20

Thr Asp Met Val Val Trp Val Thr Gly Ala Ser Ser Gly Ile Gly Glu

Glu Leu Ala Tyr Gln Leu Ser Lys Leu Gly Xaa Ser Leu Val Leu Ser 40 45 50

Ala Arg Arg Val His Glu Leu Glu Arg Val Lys Arg Arg Cys Leu Glu
55 60 65

Asn Gly Asn Leu Xaa Glu Lys Asp Ile Leu Val Leu Pro Leu Gly 70 75 80

- (2) INFORMATION FOR SEQ ID NO: 485:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 76 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -51..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 10.3 seq VSCLTLWSPGCWP/QP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 485:
- Met Thr Thr Phe Leu Pro Val Pro Gln Met Met Ala Gly Phe Ser Phe
 -50 -45 -40
- Gly Thr Phe Gly Asn Pro Pro Met Glu Ser Pro Ser Ala Trp Gln Thr
 -35
 -20
 -20
- Ile His Gln Pro Phe Ile Val Ser Cys Leu Thr Leu Trp Ser Pro Gly -15 -10 -5
- Cys Trp Pro Gln Pro Ile Gln Arg Lys Glu Trp Asp Ser Gly Thr Phe $1 \hspace{1cm} 5 \hspace{1cm} 10$
- Glu Asn Leu Arg Val Leu Ser Cys Ala Met Val Glu 15 20 25
- (2) INFORMATION FOR SEQ ID NO: 486:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 129 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate

- (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -28..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 9.5

seq LVXFSLLATAILG/AV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 486:

Met Ala Ser Lys Gly Met Arg His Phe Cys Leu Ile Ser Glu Gln Leu
-25 -20 -15

Val Xaa Phe Ser Leu Leu Ala Thr Ala Ile Leu Gly Ala Val Ser Trp
-10 -5 1

Gln Pro Thr Asn Gly Ile Phe Leu Ser Met Phe Leu Ile Val Leu Pro 5 10 15 20

Leu Glu Ser Met Ala His Gly Leu Phe His Glu Leu Gly Asn Cys Leu 25 30 35

Gly Gly Thr Ser Val Gly Tyr Ala Ile Val Ile Pro Thr Asn Phe Cys
40 45 50

Ser Pro Asp Gly Gln Pro Thr Leu Leu Pro Pro Glu His Val Gln Glu
55 60 65

Leu Asn Leu Arg Ser Thr Gly Met Leu Asn Ala Ile Gln Arg Phe Phe 70 75 80

Ala Tyr His Met Ile Glu Thr Tyr Gly Cys Asp Tyr Ser Thr Ser Gly 85 90 95 100

Leu

- (2) INFORMATION FOR SEQ ID NO: 487:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 73 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 9.3

seq VLPVILLLGAHP/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 487:

Met Ala Ala Ala Ala Trp Leu Gln Val Leu Pro Val Ile Leu Leu Leu -20 -15 -10

Leu Gly Ala His Pro Ser Pro Leu Ser Phe Phe Ser Ala Gly Pro Ala -5 5 10

Thr Val Ala Ala Ala Asp Arg Ser Lys Trp His Xaa Pro Ile Pro Ser 15 20 25

Gly Lys Asn Tyr Phe Ser Phe Gly Lys Ile Leu Phe Arg Asn Thr Thr 30 35 40

Ile Phe Leu Lys Phe Asp Gly Glu Arg
45 50

(2) INFORMATION FOR SEQ ID NO: 488:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -109..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 9.1 seq LVLAVLFFHQLVG/DP
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 488:

Met Ala Ser Pro Arg Thr Val Thr Ile Val Ala Leu Ser Val Ala Leu
-105 -100 -95

Gly Leu Phe Phe Val Phe Met Gly Thr Ile Lys Leu Thr Pro Arg Leu
-90 -85 -80

Ser Lys Asp Ala Tyr Ser Glu Met Lys Arg Ala Xaa Lys Ser Tyr Val

Arg Ala Leu Pro Leu Leu Lys Lys Met Gly Ile Asn Ser Ile Leu Leu -60 -55 -50

Arg Lys Ser Ile Gly Ala Leu Glu Val Ala Cys Gly Ile Val Met Thr
-45 -30 -35

Leu Val Pro Gly Arg Pro Lys Asp Val Ala Asn Phe Phe Leu Leu Leu -25 -15

Leu Val Leu Ala Val Leu Phe Phe His Gln Leu Val Gly Asp Pro Leu -10 -5 1

Lys

(2) INFORMATION FOR SEQ ID NO: 489:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 45 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -38..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.8

seq LLLLCALHSHIYC/IK

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 489:

Met Pro Asn Leu Ser Phe Gly Gly Leu Asp Thr Asn Gln Met Arg Val -35 -30 -25

Asn Phe Leu Ser Val Asp Val Cys Lys Leu Leu Leu Cys Ala Leu
-20 -15 -10

His Ser His Ile Tyr Cys Ile Lys Gln Ser Ala Leu Arg -5 1 5

- (2) INFORMATION FOR SEQ ID NO: 490:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 71 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -55..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.8

seq XXLLLLNVGQLLA/QT

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 490:

466

Met Gly Pro Pro Met Leu Gln Glu Ile Ser Asn Leu Phe Leu Ile Leu -55 -50 -45 -45

Leu Met Met Gly Ala Ile Phe Thr Leu Ala Ala Leu Lys Glu Ser Leu -35 -30 -25

Ser Thr Cys Ile Pro Ala Ile Val Cys Leu Xaa Xaa Leu Leu Leu Leu -20 -15 -10

Asn Val Gly Gln Leu Leu Ala Gln Thr Lys Lys Val Val Arg Pro Thr -5 1 5

Arg Lys Lys Thr Leu Ser Thr 10 15

- (2) INFORMATION FOR SEQ ID NO: 491:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 74 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -71..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.6 seq VVXFLLLLAXLIA/TY
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 491:

Met Xaa Xaa Phe Thr Asp Pro Ser Ser Val Asn Glu Lys Lys Arg Arg -70 -65 -60

Glu Arg Glu Glu Arg Gln Asn Ile Val Leu Trp Arg Gln Pro Leu Ile
-55 -50 -45 -45

Thr Leu Gln Tyr Phe Ser Leu Glu Ile Leu Val Ile Leu Lys Glu Trp

Thr Ser Lys Leu Trp His Arg Xaa Xaa Ile Val Val Xaa Phe Leu Leu -20 -15 -10

Leu Leu Ala Xaa Leu Ile Ala Thr Tyr Tyr
-5

- (2) INFORMATION FOR SEQ ID NO: 492:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 21 amino acids

- (B) TYPE: AMINO ACID
 (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.6

seq LLRGLLWXQVLCA/GP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 492:

Met Pro Leu Leu Arg Gly Leu Leu Trp Xaa Gln Val Leu Cys Ala Gly
-15 -5 1

Pro Leu His Thr Glu 5

- (2) INFORMATION FOR SEQ ID NO: 493:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 65 amino acids
 - (3) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -20..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.4

seq AVVGCLLVPPAEA/NK

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 493:

Met Lys Leu Leu Ser Leu Val Ala Val Val Gly Cys Leu Leu Val Pro
-20 -15 -10 -5

Pro Ala Glu Ala Asn Lys Ser Ser Glu Asp Ile Xaa Cys Lys Cys Ile 1 5 10

Cys Pro Pro Tyr Arg Asn Ile Ser Gly His Ile Tyr Asn Gln Asn Val

Ser Gln Lys Asp Cys Asn Cys Leu His Val Val Glu Pro Met Pro Val 30 40

Pro 45

- (2) INFORMATION FOR SEQ ID NO: 494:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 69 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -24..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.9

seq LLLPRVLLTMASG/SP

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 494:
- Met Pro Ala Leu Leu Pro Val Ala Ser Arg Leu Leu Leu Leu Pro Arg
- Val Leu Leu Thr Met Ala Ser Gly Ser Pro Pro Thr Gln Pro Ser Pro
 -5 1 5
- Ala Ser Asp Ser Gly Ser Gly Tyr Val Pro Gly Ser Val Ser Ala Ala 10 15 20
- Phe Val Thr Cys Pro Asn Glu Lys Val Ala Lys Glu Ile Ala Arg Ala 25 30 35

Val Gly Glu Lys Arg

- (2) INFORMATION FOR SEQ ID NO: 495:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 119 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide

WO 99/06550 PCT/IB98/01232

- (B) LOCATION: -108..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.9

seq LLGLLSAEQLAEA/SV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 495:

Met Cys Leu Leu Gly Ala Thr Gly Val Gly Lys Thr Leu Leu Val -105 -100 -95

Lys Arg Leu Gln Glu Val Ser Ser Arg Asp Gly Lys Gly Asp Leu Gly
-90 -85 -80

Glu Pro Pro Pro Thr Arg Pro Thr Val Gly Thr Asn Leu Thr Asp Ile -75 -70 -65

Val Ala Gln Arg Lys Ile Thr Ile Arg Glu Leu Gly Gly Cys Met Gly -60 -55 -50 -45

Pro Ile Trp Ser Ser Tyr Tyr Gly Asn Cys Arg Ser Leu Leu Phe Val

Met Asp Ala Ser Asp Pro Thr Gln Leu Ser Ala Xaa Xaa Val Gln Leu -25 -20 -15

Leu Gly Leu Leu Ser Ala Glu Gln Leu Ala Glu Ala Ser Val Leu Ile -10 -5 1

Leu Phe Asn Lys Ile Asp Asn 5 10

- (2) INFORMATION FOR SEQ ID NO: 496:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 54 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -41..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.7

seq LLCLGQLHHPGLG/RV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 496:

Met Glu Leu Pro Ala Val Asn Leu Glu Ser Asp Ser Pro Arg Ser Leu
-40 -35 -30

Ala Ala Asp Asn Leu Gly Leu His Cys Ile Leu Arg Leu Leu Cys Leu

WO 99/06550 PCT/IB98/01232

~15

-25 -20

Gly Gln Leu His His Pro Gly Leu Gly Arg Val Gly Cys Gly Ser Ala

Gly Leu His Arg Arg Arg 10

- (2) INFORMATION FOR SEQ ID NO: 497:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 53 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -51..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.6 seq PALILLFALGSLG/SG
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 497:

Met Ala Phe Leu Arg Lys Val Tyr Ser Ile Leu Ser Leu Gln Val Leu -50 -45 -40

Leu Thr Thr Val Thr Ser Thr Val Phe Leu Tyr Phe Glu Ser Val Arg -35 -25 -20

Thr Phe Val His Glu Ser Pro Ala Leu Ile Leu Leu Phe Ala Leu Gly

Ser Leu Gly Ser Gly

- (2) INFORMATION FOR SEQ ID NO: 498:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 61 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: -29..-1
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 7.6

seq PTLAIALAANAWA/FV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 498:

Met Tyr Thr Tyr Gly Asn Lys Gln His Asn Ser Pro Thr Trp Asp Asp -25 -20 -15

Pro Thr Leu Ala Ile Ala Leu Ala Ala Asn Ala Trp Ala Phe Val Leu
-10 -5 1

Phe Tyr Val Ile Pro Glu Val Ser Gln Val Thr Lys Ser Ser Pro Glu 5 15

Gln Ser Tyr Gln Gly Asp Met Tyr Pro Thr Arg Asp Leu 20 25 30

- (2) INFORMATION FOR SEQ ID NO: 499:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 42 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -32..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.6

seq WILVLALPLTVWP/WL

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 499:
- Met Gln Gln Ile Phe Ile Gln Gln Cys Arg Glu Leu Asn Phe Trp Ser

Arg Glu Pro Trp Ile Leu Val Leu Ala Leu Pro Leu Thr Val Trp Pro
-15 -5

Trp Leu Ser Pro Glu Ala Gln Pro Pro Leu
1 5 10

- (2) INFORMATION FOR SEQ ID NO: 500:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 88 amino acids

PCT/IB98/01232

- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.5

seq AVLLALLMAGLAL/OP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 500:

Met Lys Ala Val Leu Leu Ala Leu Leu Met Ala Gly Leu Ala Leu Gln -15 -5 1

Pro Gly Thr Ala Leu Leu Cys Tyr Ser Trp Xaa Ala Gln Val Xaa Asn 5 10 15

Glu Asp Cys Leu Gln Val Glu Asn Cys Thr Gln Leu Gly Glu Gln Cys
20 25 30

Trp Thr Ala Arg Ile Arg Ala Val Gly Leu Leu Thr Val Ile Ser Lys 35 40 45

Gly Cys Ser Leu Asn Cys Val Asp Xaa Ser Gln Asp Tyr Tyr Val Gly
50 60 65

Lys Lys Asn Ile Thr Cys Cys Asp

- (2) INFORMATION FOR SEQ ID NO: 501:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 82 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -16..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.1

seq QACLLGLFALILS/GK

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 501:

Met Gly Leu Gln Ala Cys Leu Leu Gly Leu Phe Ala Leu Ile Leu Ser
-15 -10 -5

Gly Lys Cys Ser Tyr Ser Pro Glu Pro Asp Gln Arg Arg Thr Leu Pro 1 5 10

Pro Gly Trp Val Ser Leu Gly Arg Ala Asp Pro Glu Glu Glu Leu Ser 20 25 30

Leu Thr Phe Ala Leu Arg Gln Gln Asn Val Glu Arg Leu Ser Glu Leu 35 40 45

Val Gln Ala Val Ser Asp Pro Ser Ser Pro Gln Tyr Gly Lys Tyr Leu 50 60

Thr Arg 65

(2) INFORMATION FOR SEQ ID NO: 502:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 127 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -29..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7

seq LGSGLGLSPGTSS/GR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 502:

Met Arg Pro Gly Gln Val Ser Leu Leu Gly Pro Asp Ala Val Ser Val -25 -20 -15

Leu Gly Ser Gly Leu Gly Leu Ser Pro Gly Thr Ser Ser Gly Arg Asn -10 -5 1

Pro Asp Pro Gly Ser Gly Pro Gly Thr Leu Pro Xaa Xaa Ser Xaa Gln 5

Asn Pro Ser Pro Ala Pro Asp Pro Pro Pro Ala Leu Leu Trp Asn 20 25 30 35

Leu Leu Thr Gln Arg Leu Gly Thr Thr Leu Val Pro Thr Leu Cys Pro
40 45 50

Ala Gin Thr Leu Ile Leu Cys Pro Ala Gin Thr Leu Ile Leu Cys Pro
55 60 65

Xaa Leu Ile Pro Thr Leu Cys Pro Ala Leu Xaa Pro Val Leu Pro Xaa 70 75 80

Val Ala Leu Ser Ala Gln Pro Ser Leu Pro Ala Arg Val Gln Ser 85 90 95

- (2) INFORMATION FOR SEQ ID NO: 503:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 43 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -33..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.8 seq FTSASLLLPMSTG/MP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 503:
- Met Ile Asn Pro Ser Val Pro Ser Lys Ser Asn Ser His Pro Phe Leu
 -30 -25 -20
- Ser Thr Val Met Phe Thr Ser Ala Ser Leu Leu Leu Pro Met Ser Thr -15 -10 -5
- Gly Met Pro Thr Gln Asn Cys Phe Thr Pro Lys
 1 5 10
- (2) INFORMATION FOR SEQ ID NO: 504:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 108 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -68..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.7

seq IACLAWWIGGGSG/XN

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 504:
- Met Ser Glu Lys Glu Xaa Asn Phe Pro Pro Leu Pro Lys Phe Ile Pro
 -65 -60 -55
- Val Lys Pro Cys Phe Tyr Gln Asn Phe Ser Asp Glu Ile Pro Val Glu -50 -45 -40
- His Gln Val Leu Val Lys Arg Ile Tyr Arg Leu Trp Met Phe Tyr Cys -35 -30 -25
- Ala Thr Leu Gly Val Asn Leu Ile Ala Cys Leu Ala Trp Trp Ile Gly
 -20 -15 -10 -5
- Gly Gly Ser Gly Xaa Asn Phe Gly Leu Ala Phe Val Trp Leu Leu Leu $1 \hspace{1.5cm} 5 \hspace{1.5cm} 10$
- Phe Thr Pro Cys Gly Tyr Val Cys Trp Phe Arg Pro Val Tyr Lys Ala 15 20 25
- Phe Arg Ala Asp Ser Ser Phe Asn Phe Met Ala Leu $30 \hspace{1cm} 35 \hspace{1cm} 40$
- (2) INFORMATION FOR SEQ ID NO: 505:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 63 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.7 seq ILRLYFFLQLAHS/GY
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 505:
- Met Asn Pro Thr Lys Leu Ile Leu Lys Thr Ile Leu Arg Leu Tyr Phe
 -20
 -15
 -10
- Phe Leu Gln Leu Ala His Ser Gly Tyr Thr Lys Leu Gln Lys Lys Tyr
 -5
- Met Lys Ser Arg Tyr Glu Gl
n Val Asp Leu Val Gly Lys Met Xaa Gl
n 10 20 25
- Lys Ala Ala Thr Thr Val Xaa His Leu Ala Ile Gln Cys His Trp

35

30

40

(2) INFORMATION FOR SEQ ID NO: 506:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 123 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.7

seq SXXCFVSVPPASA/IP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 506:

Met Ala Ser Ser Ser Pro Asp Ser Pro Cys Ser Xaa Xaa Cys Phe Val

Ser Val Pro Pro Ala Ser Ala Ile Pro Xaa Val Xaa Xaa Ala Xaa Asn -5 l 5

Ser Asp Xaa Pro Arg Asp Glu Val Gln Glu Val Val Phe Val Pro Ala 10 15 20 25

Gly Thr His Thr Pro Gly Ser Arg Leu Gln Cys Thr Tyr Ile Glu Val $30 \hspace{1cm} 35 \hspace{1cm} 40$

Glu Gln Val Ser Lys Thr His Ala Val Ile Leu Ser Arg Pro Ser Trp
45 50 55

Leu Trp Gly Ala Glu Met Gly Xaa Thr Ser Met Val Ser Ala Leu Ala 60 65 70

Thr Arg Leu Cys Gly Arg Arg Ser Gln Leu Gly Arg Ala Xaa Ala Leu 75 80 85

Leu Gly Met Asp Leu Leu Arg Cys Arg Pro Cys 90 95 100

(2) INFORMATION FOR SEQ ID NO: 507:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 46 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -39..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.7

seq XLIAXLEPPGAMA/VR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 507:

Met Xaa Pro Val Leu Ala Ala Leu Ala His Val Leu Cys Pro Tyr Met
-35 -30 -25

Ala Pro Gly Leu Cys Arg Glu Pro Ile Arg Xaa Leu Ile Ala Xaa Leu
-20 -15 -10

Glu Pro Pro Gly Ala Met Ala Val Arg Arg Leu Pro Ser Ala

- (2) INFORMATION FOR SEQ ID NO: 508:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 85 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -45..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.7

seq PMLGLAAFRWIWS/RE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 508:

Met Asn Asn Leu Asn Asp Pro Pro Asn Trp Asn Ile Arg Pro Asn Ser
-45 -35 -30

Arg Ala Asp Gly Gly Asp Gly Ser Arg Trp Asn Tyr Ala Leu Leu Val
-25 -20 -15

Fro Met Leu Gly Leu Ala Ala Phe Arg Trp Ile Trp Ser Arg Glu Ser
-10 -5 1

Gim Lys Glu Val Glu Lys Glu Arg Glu Ala Tyr Arg Arg Thr Ala 5 10 15 Ala Phe Gln Gln Asp Leu Glu Ala Lys Tyr His Ala Met Ile Ser Xaa 20 25 30

Asn Arg Arg Ala Val

- (2) INFORMATION FOR SEQ ID NO: 509:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 29 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -19..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.6 seq AALCSLFFFLSLQ/EI
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 509:

Met Leu Leu Leu Phe Leu Ala Ala Leu Cys Ser Leu Phe Phe Phe Leu
-15 -10 -5

Ser Leu Gln Glu Ile Ala Pro Gln Asp Pro Lys Pro Gly 1 5 10

- (2) INFORMATION FOR SEQ ID NO: 510:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 59 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -47..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.5

seq IIVCLFAFLVAHC/FL

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 510:

Met Leu Phe Leu Gly Lys Val Leu Ile Val Cys Ser Thr Gly Leu Ala

Gly Ile Met Leu Leu Asn Tyr Gln Gln Asp Tyr Thr Val Trp Val Leu
-30 -25 -20

Pro Leu Ile Ile Val Cys Leu Phe Ala Phe Leu Val Ala His Cys Phe -15 -5 1

Leu Ser Ile Tyr Glu Met Val Val Asp Ala Arg
5 10

- (2) INFORMATION FOR SEQ ID NO: 511:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 43 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (3) LOCATION: -38..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.3 seq LLLLVHSFWFTVC/TP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 511:

Met Gln Gly Ile Pro Ile Leu Thr Pro Val Thr Thr Gln Ser Ile Ala
-35
-30
-25

Ile Ser Ile Val Leu Thr Val Glm Gly Leu Leu Leu Val His Ser
-20 -15 -10

Phe Trp Phe Thr Val Cys Thr Pro Val Val Phe
-5 1 5

- (2) INFORMATION FOR SEQ ID NO: 512:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 54 amino acids
 - (E) TYPE: AMINO ACID
 - (C) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -27..-1
 - (C) PDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.3

seq LFCVLLSLRPHTS/GT

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 512:
- Met Gln Asn Phe Cys His His Leu Ala Ile Cys Thr Val Ile Leu Phe
 -25
 -20
 -15
- Cys Val Leu Leu Ser Leu Arg Pro His Thr Ser Gly Thr Leu Trp Ala . -10 -5 1 5
- Ser Ser Ala His Gly Leu His Leu Ala Pro Ala Glu Pro Gln Leu Ser 10 15 20

Cys Trp Met Cys Cys Ala 25

- (2) INFORMATION FOR SEQ ID NO: 513:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 135 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -64..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.3

seq VLMRLVASAYSIA/QK

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 513:
- Gly His Xaa Xaa Gly Xaa Gly Ser Tyr Asp Xaa Ala Leu Xaa Leu Leu -45 -40 -35
- Leu Lys Cys Leu Trp Ser Asn Val Val Pro Glu Cys Thr Met Ala Ser
- Ser Asn Thr Val Leu Met Arg Leu Val Ala Ser Ala Tyr Ser Ile Ala
 -15
 -10
 -5
- Gln Lys Ala Gly Met Ile Val Arg Arg Val Ile Ala Glu Gly Asp Leu

1 5 10 15

Gly Ile Val Glu Lys Thr Cys Ala Thr Asp Leu Gln Thr Lys Ala Asp
20 25 30

Arg Leu Ala Gln Met Ser Ile Cys Ser Ser Leu Xaa Xaa Lys Phe Pro 35 40 45

Lys Leu Xaa Ile Ile Gly Glu Glu Asp Leu Pro Ser Glu Glu Val Asp
50 60

Gln Glu Leu Ile Glu Asp Xaa 65 70

- (2) INFORMATION FOR SEQ ID NO: 514:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 28 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.2 seq LEMLXAFASHIXA/RD
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 514:

Met Arg Gly Ala His Leu Thr Ala Leu Glu Met Leu Xaa Ala Phe Ala -20 -15 -10

Ser His Ile Xaa Ala Arg Asp Ala Ala Gly Ser Gly

- (2) INFORMATION FOR SEO ID NO: 515:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 141 amino acids
 - (3) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:

WO 99/06550 PCT/IB98/01232 482

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: -139..-1
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 6.2

seq FGLLHQLSQCVTS/LE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 515:

Met Glu Val Gly Leu Pro Ala Ile Thr Leu Phe Leu Thr Ser Ala Ser -130

Ser Pro Val Val Ala Thr Thr Met Asp Gln Glu Pro Val Gly Gly Val

Glu Arg Gly Glu Ala Val Ala Ala Ser Gly Xaa Ala Ala Ala Ala Ala -100

Phe Gly Glu Ser Ala Gly Gln Met Ser Asn Glu Arg Gly Phe Glu Asn

Val Glu Leu Gly Val Ile Gly Lys Lys Lys Lys Val Pro Arg Arg Val

Ile His Phe Val Ser Gly Glu Thr Met Glu Glu Tyr Ser Thr Asp Glu

Asp Xaa Val Asp Gly Leu Glu Lys Xaa Met Phe Cys Leu Leu Ile -35

Arg Gln Asn Leu Pro Gly Val Pro Thr Tyr Gly Phe Thr Cys Phe Gly

Leu Leu His Gln Leu Ser Gln Cys Val Thr Ser Leu Glu -5

- (2) INFORMATION FOR SEQ ID NO: 516:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 57 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -43..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.1 seq SAATLASLGGTSS/RR
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 516:

Met Lys Glu Leu Glu Arg Gln Gln Lys Glu Val Glu Glu Arg Pro Glu
-40 -35 -30

Lys Asp Phe Thr Glu Lys Gly Ser Arg Asn Met Pro Gly Leu Ser Ala
-25 -20 -15

Ala Thr Leu Ala Ser Leu Gly Gly Thr Ser Ser Arg Arg Gly Ser Gly
-10 -5 1 5

Asp Thr Ser Ile Ser Ile Asp Pro Glu 10

- (2) INFORMATION FOR SEQ ID NO: 517:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 92 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.1 seq VLVILCIVTVCVT/IV
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 517:

Met Ser Met Gly Phe Met Met Leu Val Leu Val Ile Leu Cys Ile Val -20 -15 -10

Thr Val Cys Val Thr Ile Val Cys Thr Tyr Phe Leu Leu Asn Ala Glu -5 - 10

Asp Tyr Arg Trp Gln Trp Thr Ser Phe Leu Ser Ala Ala Ser Thr Ala 15 20 25

Ile Tyr Val Tyr Met Tyr Ser Phe Tyr Tyr Tyr Phe Phe Lys Thr Lys 30 35 40

Met Tyr Gly Leu Phe Gln Thr Ser Phe Tyr Phe Gly Tyr Met Ala Val 45 50 55

Phe Ser Thr Ala Leu Gly Ile Met Cys Gly Ala Ile 60 65 70

- (2) INFORMATION FOR SEQ ID NO: 518:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids

(B) TYPE: AMINO ACID(D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: PROTEIN

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

(ix) FEATURE:

(A) NAME/KEY: sig_peptide

(B) LOCATION: -70..-1

(C) IDENTIFICATION METHOD: Von Heijne matrix

(D) OTHER INFORMATION: score 6

seq LLFPLTLVRSFWS/DM

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 518:

Met Met Glu Leu Xaa Leu Lys Xaa Xaa Thr Lys Xaa Glu Xaa Glu Ser -70 -65 -60 -55

Ala Cys Thr Glu Ala Tyr Ser Gln Ser Asp Glu Gln Tyr Ala Cys His
-50 -45 -40

Leu Gly Cys Gln Asn Gln Leu Pro Phe Ala Glu Leu Arg Gln Glu Gln -35 -30 -25

Leu Met Ser Leu Met Pro Lys Met His Leu Leu Phe Pro Leu Thr Leu -20 -15 -10

Val Arg Ser Phe Trp Ser Asp Met Met Asp Ser Ala Gln Ser Phe Xaa-5 1 5 10

Thr Ser Ser Trp Thr Phe Tyr Leu Gln Ala Asp Xaa Gly Xaa Ile Val 15 20 25

Ile Xaa Gl
n Ser Lys Pro Glu Ile Gl
n Tyr Ala Pro His Leu Glu Gl
n 30 35 40

Glu

- (2) INFORMATION FOR SEQ ID NO: 519:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 27 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -24..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix

. (D) OTHER INFORMATION: score 6

seq GLILLFASHLINO/FS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 519:

Met Val Ser Asn Ala Ser Glu Thr Ser Cys Leu Gly Leu Ile Leu Leu -20 -15 -10

Phe Ala Ser His Leu Ile Asn Gln Phe Ser Ser -5

- (2) INFORMATION FOR SEQ ID NO: 520:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 124 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -73..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6 seq LIVFISVCTALLA/EG
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 520:

Met Pro Arg Lys Arg Lys Cys Asp Leu Arg Ala Val Arg Val Gly Leu
-70 -65 -60

Leu Leu Gly Gly Gly Val Tyr Gly Ser Arg Phe Arg Phe Thr Phe
--55 -50 -45

Pro Gly Cys Arg Ala Leu Ser Pro Trp Arg Val Arg Xaa Gln Arg Arg
-40 -35 -30

Arg Cys Glu Met Ser Thr Met Phe Ala Asp Thr Leu Leu Ile Val Phe -25 -15 -10

Ile Ser Val Cys Thr Ala Leu Leu Ala Glu Gly Ile Thr Trp Val Leu
-5 5

Val Tyr Arg Thr Asp Lys Tyr Lys Arg Leu Lys Ala Glu Val Glu Lys
10 15 20

Gln Ser Lys Lys Tyr Leu Met Val Glu Trp Trp Gln Xaa Phe Leu Phe $25 \hspace{1cm} 30 \hspace{1cm} 35$

Tyr Pro Ser Phe Leu Xaa Pro Lys Xaa Val Ser Ser 40 45 50

- (2) INFORMATION FOR SEQ ID NO: 521:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 106 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq LGAAALALLLANT/DV

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 521:
- Met Gly Met Trp Ser Ile Gly Ala Gly Ala Leu Gly Ala Ala Ala Leu
 -20 -15 -10
- Ala Leu Leu Leu Ala Asn Thr Asp Val Phe Leu Ser Lys Pro Gln Lys
 -5 1 5
- Ala Ala Leu Glu Tyr Leu Glu Asp Ile Asp Leu Lys Thr Leu Glu Lys 10 20 25
- Glu Pro Arg Thr Phe Lys Ala Lys Glu Leu Trp Glu Lys Asn Gly Ala 30 35 40
- Val Ile Met Ala Val Arg Arg Pro Gly Cys Phe Leu Cys Arg Glu Glu 45 50 55
- Ala Ala Asp Leu Ser Ser Leu Lys Ser Met Leu Asp Gln Leu Gly Val 60 65 70
- Pro Leu Tyr Ala Val Val Lys Glu Gln Arg
- (2) INFORMATION FOR SEQ ID NO: 522:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 58 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -31..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.9

seq LPLLLVANAGTAA/VG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 522:

Met Asp Val Ala Phe Leu Glu Xaa Leu Ile Lys Asp Asp Ile Glu Arg
-30 -25 -20

Gly Arg Leu Pro Leu Leu Val Ala Asn Ala Gly Thr Ala Ala Val -15 -10 -5 1

Gly His Thr Asp Lys Ile Gly Arg Leu Lys Glu Leu Cys Glu Gln Tyr
5 10 15

Gly Ile Trp Leu His Val Glu Gly Val Asn 20 25

- (2) INFORMATION FOR SEQ ID NO: 523:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 18 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -16..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.8 seq LFNLLWLALACSP/VW
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 523:

Met Arg Thr Leu Phe Asn Leu Leu Trp Leu Ala Leu Ala Cys Ser Pro -15 -10 -5

Val Trp

- (2) INFORMATION FOR SEQ ID NO: 524:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 39 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: PROTEIN

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -33..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.8

seq FICLQWALPHSEA/GD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 524:

Met Asn Ala Gln Pro Gly Leu Xaa Leu Asp Cys Ile Thr Arg Phe Leu
-30 -25 -20

Thr Xaa Gly Gln Phe Ile Cys Leu Gln Trp Ala Leu Pro His Ser Glu -15 -10 -5

Ala Gly Asp Phe Glu Ala Lys
1 5

- (2) INFORMATION FOR SEQ ID NO: 525:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 78 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -69..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.7 seq LCRLLCLVRLFCC/SS
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 525:

Met Gly Lys Glu Trp Gly Trp Gln Glu Met Glu Asn Gly Gly Ala Ala
-65 -60 -55

Pro Ala Trp Gly Ala Gly Pro Pro Val His Pro Ala Pro Pro Pro Val -50 -45 -40

Glu Lys Thr Leu Ser Trp Gly Cys Gly Phe Gly Leu His Ser Gly Phe
-35 -30 -25

Gly Gly Ser Gly Gly Gly Val Gly Leu Cys Arg Leu Leu Cys Leu Val

-20

-15

Arg Leu Phe Cys Cys Ser Ser Ile Leu Tyr Gln Arg Gln Lys 1

- (2) INFORMATION FOR SEQ ID NO: 526:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 33 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -29..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.7

seq AALLLTATVRLSA/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 526:

Met Ala Ala Pro Ser Gly Gly Trp Asn Gly Val Gly Ala Ser Leu Trp

Ala Ala Leu Leu Thr Ala Thr Val Arg Leu Ser Ala Ser Pro Gly

Pro

- (2) INFORMATION FOR SEQ ID NO: 527:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 130 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -48..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.7 seq LLLFFGKLLVVGG/VG
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 527:

Met Ile Ala Ile Tyr Gly Lys Asn Phe Cys Val Ser Ala Lys Asn Ala
-45 -40 -35

Phe Met Leu Leu Met Arg Asn Ile Val Arg Val Val Leu Asp Lys -30 -25 -20

Val Thr Asp Leu Leu Phe Phe Gly Lys Leu Leu Val Val Gly Gly
-15 -5

Val Gly Val Leu Ser Phe Phe Phe Phe Ser Gly Arg Ile Pro Gly Leu 1 5 10 15

Gly Lys Asp Phe Lys Ser Pro His Leu Asn Tyr Tyr Trp Leu Pro Xaa 20 25 30

Met Thr Ser Ile Leu Gly Ala Tyr Val Ile Ala Ser Gly Phe Phe Ser 35 . 40 . 45

Val Phe Gly Met Cys Val Asp Thr Leu Phe Leu Cys Phe Leu Glu Asp 50 55 60

Leu Glu Arg Thr Thr Ala Pro Trp Thr Ala Leu Leu His Val Gln Glu 65 70 75 80

Leu Leu

- (2) INFORMATION FOR SEQ ID NO: 528:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 96 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -91..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.7 seq SVLELIVASVCQS/HI
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 528:

Met Glu Arg Asn Cys Lys Gly Ser Phe Gly Val Ile Lys Glu Gly Asp
-90 -85 -80

Thr Asp Thr Xaa Glu Thr Lys Ala Arg Arg Thr Val Trp Glu Pro Arg -75 -65 -60

Gly Arg Tyr Ser Phe Arg Xaa Thr Pro Arg Pro Ala Tyr Pro Val Glu
-55 -50 -45

Gln Cys Gly Phe Ala Arg Arg Ala Leu Glu Leu Glu Ile Arg Lys
-40 -35 -30

His Ser Pro Glu Val Cys Glu Pro Pro Asn Ile Pro Val Thr Ser Val -25 -20 -15

Leu Glu Leu Ile Val Ala Ser Val Cys Gln Ser His Ile Arg Thr Thr
-10 -5 1 5

(2) INFORMATION FOR SEQ ID NO: 529:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 93 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -66..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.7 seq LYMLAEALPVSHG/AH
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 529:

Met Phe Val Glu Tyr Arg Lys Gln Leu Lys Leu Leu Leu Asp Arg Leu
-65 -55

Ala Gln Val Ser Pro Glu Leu Leu Leu Ala Ser Val Arg Arg Val Phe
-50
-45
-40
-35

Ser Ser Thr Leu Gln Asn Trp Gln Thr Thr Arg Phe Met Glu Val Glu -30 -25 -20

Val Ala Ile Arg Leu Leu Tyr Met Leu Ala Glu Ala Leu Pro Val Ser
-15 -10 -5

His Gly Ala His Phe Ser Gly Asp Val Ser Lys Ala Ser Ala Leu Gln $1 \hspace{1.5cm} 5 \hspace{1.5cm} 10$

Asp Met Met Arg Thr Leu Val Thr Ser Gly Val Ser Gly 15 20 25

(2) INFORMATION FOR SEQ ID NO: 530:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR

(ii) MOLECULE TYPE: PROTEIN

(vi) ORIGINAL SOURCE:

- (A) ORGANISM: Homo Sapiens
- (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.7

seq IIFLIQWHGSVFO/EF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 530:

Met Leu Leu Gly Thr Ser Asn Ile Ile Ile Phe Leu Ile Gln Trp His -20 -15 -10

Gly Ser Val Phe Gln Glu Phe
-5

- (2) INFORMATION FOR SEQ ID NO: 531:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 54 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -20..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.6

seq AFVXACVLSLIST/IY

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 531:

Met Xaa Asn Arg Phe Ala Thr Ala Phe Val Xaa Ala Cys Val Leu Ser -20 -15 -10 -5

Leu Ile Ser Thr Ile Tyr Met Ala Ala Ser Ile Gly Thr Asp Phe Trp $1 \hspace{1cm} 5 \hspace{1cm} 10$

Tyr Glu Tyr Arg Ser Pro Val Gln Glu Asn Ser Ser Asp Leu Asn Lys 15 20 25

Ser Ile Tro Asp Glu Leu 30

(2) INFORMATION FOR SEO ID NO: 532:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 19 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -13..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.6

seq MSLTSGFLRVSQG/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 532:

Met Ser Leu Thr Ser Gly Phe Leu Arg Val Ser Gln Gly Ser Pro Asn -10 -5 1

Leu Ser Gln 5

- (2) INFORMATION FOR SEQ ID NO: 533:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 86 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -63..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.6

seq AIRTLFSVTGILA/EQ

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 533:

Met Ala Asn Pne Lys Gly His Ala Leu Pro Gly Ser Phe Phe Leu Ile
-60 -55 -50

Ile Gly Leu Cys Trp Ser Val Lys Tyr Pro Leu Lys Tyr Phe Ser His

494

Thr Arg Lys Asn Ser Pro Leu His Tyr Tyr Gln Arg Leu Glu Ile Val

Glu Ala Ala Ile Arg Thr Leu Phe Ser Val Thr Gly Ile Leu Ala Glu -15 -10 -5 1

Gln Phe Val Pro Asp Gly Pro His Leu His Leu Tyr His Glu Asn His 5 10 15

Trp Ile Lys Leu Met Asn 20

- (2) INFORMATION FOR SEQ ID NO: 534:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 73 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -52..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.5 seq AGLLFGSLAGLGA/YQ
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 534:

Met Gln Asp Thr Gly Ser Val Val Pro Leu His Trp Phe Gly Phe Gly -50 -45 -40

Tyr Ala Ala Leu Val Ala Ser Gly Gly Ile Ile Gly Tyr Val Lys Ala
-35 - -30 -25

Gly Ser Val Pro Ser Leu Ala Ala Gly Leu Leu Phe Gly Ser Leu Ala -20 -15 -10 -5

Gly Leu Gly Ala Tyr Gln Leu Ser Gln Asp Pro Arg Asn Val Trp Val

Phe Leu Ala Thr Ser Gly Thr Leu Ala 15 20

- (2) INFORMATION FOR SEQ ID NO: 535:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 47 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR

- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig peptide
 - (B) LOCATION: -35..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.4

seq CCALLTSLXCIWG/PA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 535:

Met Glu Xaa Gly Leu Lys Ser Ala Asp Pro Arg Asp Gly Thr Gly Tyr
-35
-20
-20

Thr Xaa Xaa Xaa Tyr Cys Cys Ala Leu Leu Thr Ser Leu Xaa Cys -15 -10 -5

Ile Trp Gly Pro Ala Tyr Leu Gln Leu Ala His Gly Tyr Val Lys 1 5 10

- (2) INFORMATION FOR SEQ ID NO: 536:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 77 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -42..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.4

seq ITGVILLAVGIWG/KV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 536:

Met Ala Ser Pro Ser Arg Arg Leu Gln Thr Lys Pro Val Ile Thr Cys
-40 -35 -30

Phe Lys Ser Val Leu Leu Ile Tyr Thr Phe Ile Phe Trp Ile Thr Gly -25 -20 -15

Val Ile Leu Leu Ala Val Gly Ile Trp Gly Lys Val Ser Leu Glu Asn -10 5 1 5

Tyr Phe Ser Leu Leu Asn Glu Lys Ala Thr Asn Val Pro Phe Val Leu 10 15 20 Ile Ala Thr Gly Thr Val Ile Ile Leu Leu Gly Thr Leu 25 30 35

- (2) INFORMATION FOR SEQ ID NO: 537:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 89 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -67..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.2 seq LSVSLLPCAGAWS/LL
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 537:
- Met Phe Ser Arg Glu Leu Ala Pro Thr Arg Ile Gly Gly Ala Ser Ser
 -65 -60 -55
- Gly Ser Arg Ser Gly Gly Thr Leu Ile Ser Thr Ala Pro Leu Thr Thr -50 -45 -40
- Arg Val Leu Asn Pro Thr Ala Gln Cys Phe Cys Leu Asp Cys Thr Leu
 -35 -30 -25 -20
- Arg Arg Met Gln Thr His Leu Ser Val Ser Leu Leu Pro Cys Ala Gly -15 -10 -5
- Ala Trp Ser Leu Leu Xaa Ser Lys Lys Val Ile Leu Pro Ser Cys Ser 1 5 10
- Ser Ile Leu Xaa Thr Val Val Val Ile 15 20
- (2) INFORMATION FOR SEQ ID NO: 538:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 54 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -29..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.1

seq LLMLGVTLPNSYW/RV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 538:

Met Ser Met Ala Val Glu Thr Phe Gly Phe Phe Met Ala Thr Val Gly -25 -20 -15

Leu Leu Met Leu Gly Val Thr Leu Pro Asn Ser Tyr Trp Arg Val Ser -10 -5 1

Thr Val His Gly Asn Val Ile Xaa Thr Asn Xaa Ile Phe Glu Asn Leu $5 \hspace{1.5cm} 10 \hspace{1.5cm} 15$

Trp Phe Ser Ser Ala Gly 20 25

- (2) INFORMATION FOR SEQ ID NO: 539:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 56 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -20..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5

seq XFLXLXXLSXXWP/XD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 539:

Met Glu Lys Ile Pro Val Ser Xaa Phe Leu Xaa Leu Xaa Xaa Leu Ser -20 -15 -10 -5

Thr Lys Asp Ser Arg Xaa Lys Leu Pro Gln Thr Leu Ser Arg Gly Trp

Gly Asp Gln Leu Ile Trp Thr Arg
30 35

- (2) INFORMATION FOR SEQ ID NO: 540:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 85 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -67..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5

seq LILERPLVPSAEA/SG

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 540:
- Met His Ser Ala Glu Glu Pro Leu Xaa Leu Ala Ala Leu Arg Gly Ala
 -65 -60 -55
- Arg Gly His Leu Pro Cys Gly Ser Arg His His Val Gly Ser Leu Ala -50 -45 -40
- Pro Ala Ser Val Pro Ala Pro Gly Ala Cys Leu Trp Val Cys Glu Trp
 -35 -30 -25 -20
- Glu Thr Leu Leu Pro Gly Leu Ile Leu Glu Arg Pro Leu Val Pro Ser -15 -10 -5
- Ala Glu Ala Ser Gly Ala Gly Lys Leu Ser Arg Lys Glu Ala Leu Leu

 1 5 10

Ser Asn Tyr Ala Leu 15

- (2) INFORMATION FOR SEQ ID NO: 541:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 50 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -43..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix

- (D) OTHER INFORMATION: score 4.9 seq GLWLALVDGLVRX/AP
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 541:

Met Ala Gly Gln Phe Arg Ser Tyr Val Trp Asp Pro Leu Leu Ile Leu
-40 -35 -30

Ser Gln Ile Val Leu Met Gln Thr Val Tyr Tyr Gly Ser Leu Gly Leu
-25
-20
-15

Trp Leu Ala Leu Val Asp Gly Leu Val Arg Xaa Ala Pro Arg Trp Ile -10 -5 1 5

Xaa Gly

- (2) INFORMATION FOR SEQ ID NO: 542:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 90 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -78..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.9 seq VGAVFGLTTCISA/HV
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 542:

Met Ala Pro Lys Val Phe Arg Gln Tyr Trp Asp Ile Pro Asp Gly Thr

Asp Cys His Arg Lys Ala Tyr Ser Thr Thr Ser Ile Ala Ser Val Ala
-60 -55 -50

Gly Leu Thr Ala Ala Ala Tyr Arg Val Thr Leu Asn Pro Pro Gly Thr -45 -35

Phe Leu Glu Gly Val Ala Lys Val Gly Gln Tyr Thr Phe Thr Ala Ala
-30 -25 -20 -15

Ala Val Giy Ala Val Phe Gly Leu Thr Thr Cys Ile Ser Ala His Val -10 -5 1

Arg Glu Lys Pro Asp Asp Pro Leu Asn Arg

(2) INFORMATION FOR SEQ ID NO: 543:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 22 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.9

seq WLQVLPVILLLLG/VP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 543:

Met Ala Ala Ala Trp Leu Gln Val Leu Pro Val Ile Leu Leu Leu -15 -10 -5

Leu Gly Val Pro Pro Ser

- (2) INFORMATION FOR SEQ ID NO: 544:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 41 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -37..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8

seq LLILDMNVLYTDA/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 544:

Met Glu Ile Tyr Phe Ile Phe Cys Ile Ile Val Pro Ile Ala Ala Ala -35

Thr Val Tyr Lys Ser Trp Cys Leu Leu Leu Ile Leu Asp Met Asn Val $^{-20}$ $^{-15}$ $^{-10}$

Leu Tyr Thr Asp Ala Ser Pro Leu Gly -5

- (2) INFORMATION FOR SEQ ID NO: 545:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 50 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -31..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8

seq VLLAIGMFFTAWF/FV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 545:

Met Ser Arg Tyr Thr Ser Pro Val Asn Pro Ala Val Phe Pro His Leu
-30 -25 -20

Thr Val Val Leu Leu Ala Ile Gly Met Phe Phe Thr Ala Trp Phe Phe -15 -5 1

Val Tyr Glu Val Thr Ser Thr Lys Tyr Thr Arg Asp Ile Tyr Lys Glu 5 10 15

Leu Gln

- (2) INFORMATION FOR SEQ ID NO: 546:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 38 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -35..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8

seq LMLSSSLPLLIWL/KD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 546:

Met Arg Leu Ala Ala Glu Ala His Pro Gly Arg Thr His Thr Leu Phe
-35 -20 -25

Arg Arg Leu Lys Pro Phe Leu Met Leu Ser Ser Ser Leu Pro Leu Leu -15 -10 -5

Ile Trp Leu Lys Asp Arg

- (2) INFORMATION FOR SEQ ID NO: 547:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 55 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -39..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8 seq IILFSAIVGFIYG/YV
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 547:

Met Leu Glu His Leu Xaa Ser Leu Pro Thr Gln Met Asp Tyr Lys Gly
-35 -30 -25

Gln Lys Leu Ala Xaa Gln Met Phe Gln Gly Ile Ile Leu Phe Ser Ala - -20 -15 -10

Ile Val Gly Phe Ile Tyr Gly Tyr Val Ala Glu Gln Phe Gly Trp Thr
-5 1 5

Val Tyr Ile Val Met Ala Gly 10 15

- (2) INFORMATION FOR SEQ ID NO: 548:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 19 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:

(A) ORGANISM: Homo Sapiens(F) TISSUE TYPE: Normal prostate

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -16..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.8

seq SKVLFCSFSNVLG/FD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 548:

Met Glu Tyr Ser Lys Val Leu Phe Cys Ser Phe Ser Asn Val Leu Gly -15 -5

Phe Asp Tyr

- (2) INFORMATION FOR SEQ ID NO: 549:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 60 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -26..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.7

seq LIMQLGSVLLTRC/PF

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 549:

Met Ala Ser Lys Ile Gly Ser Arg Arg Trp Met Leu Gln Leu Ile Met -25 -20 -15

Gln Leu Gly Ser Val Leu Leu Thr Arg Cys Pro Phe Trp Gly Cys Phe -10 -5 1 5

Ser Gln Leu Met Leu Tyr Ala Glu Arg Ala Glu Ala Arg Arg Lys Pro 10 15 20

Asp Ile Pro Val Pro Tyr Leu Tyr Phe Asp Ser Gly 25 30

- (2) INFORMATION FOR SEQ ID NO: 550:
 - (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 79 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -52..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.7

seq LGLALGRLEGGSA/RH

PCT/IB98/01232

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 550:

Met Glu His Tyr Arg Lys Ala Gly Ser Val Glu Leu Pro Ala Pro Ser
-50 -45 -40

Pro Met Pro Gln Leu Pro Pro Asp Thr Leu Glu Met Arg Val Arg Asp -35 -30 -25

- Gly Ser Lys Ile Arg Asn Leu Leu Gly Leu Ala Leu Gly Arg Leu Glu
 -20 -15 -10 -5
- Gly Gly Ser Ala Arg His Val Val Phe Ser Gly Ser Gly Arg Ala Ala I 5 10
- Gly Lys Ala Val Ser Cys Ala Glu Ile Val Lys Arg Arg Val Pro \$15\$ \$20\$ \$25\$
- (2) INFORMATION FOR SEQ ID NO: 551:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 73 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -26..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.6

seq LIALTCLDGTTVS/AE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 551:

Met Asn Ala Leu Met Val Leu Phe Asn Val Thr Val Val Leu Ile Ala

-25 -20 -15

Leu Thr Cys Leu Asp Gly Thr Thr Val Ser Ala Glu Met Ala Thr Met -10 -5 1

Thr Met Gly Cys Phe His Gln Val Glu Asn Arg Val Lys Ile Leu Met 10 15 20

Ser Val Gly Pro Gly Gly Thr Ala Val Pro Met Ile Pro Phe Ala Ser 25 30 35

Ile Trp Met Ala Asp Met Ile Xaa Asp

- (2) INFORMATION FOR SEQ ID NO: 552:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 49 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -45..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.6 seq VLVYLVTAERVWS/DD
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 552:

Met Asn Trp Ser Ile Phe Glu Gly Leu Leu Ser Gly Val Asn Lys Tyr
-45 -35 -30

Ser Thr Ala Phe Gly Arg Ile Trp Leu Ser Leu Val Phe Ile Phe Arg
-25 -20 -15

Val Leu Val Tyr Leu Val Thr Ala Glu Arg Val Trp Ser Asp Asp His -10 -5 1

Lys

- (2) INFORMATION FOR SEQ ID NO: 553:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 53 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -16..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.5

seq SLFIYIFXTCSNT/SP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 553:

Met Ile Ile Ser Leu Phe Ile Tyr Ile Phe Xaa Thr Cys Ser Asn Thr -15 -10 -5

Ser Pro Ser Tyr Gln Xaa Thr Gln Leu Gly Leu Gly Leu Pro Ser Ala 1 5 10

Gln Trp Trp Pro Leu Thr Gly Arg Arg Met Gln Cys Cys Arg Leu Phe
20 25 30

Cys Phe Xaa Leu Gln

- (2) INFORMATION FOR SEQ ID NO: 554:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 32 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix-) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -16..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.4

seq LNSLSALAELAVG/SR

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 554:
- Met Phe Arg Leu Asn Ser Leu Ser Ala Leu Ala Glu Leu Ala Val Gly
 -15 -5

Ser Arg Trp Tyr His Gly Gly Ser Gln Pro Ile Gln Ile Arg Arg Arg 1 5 10 15

(2) INFORMATION FOR SEQ ID NO: 555:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 44 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -17..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.4

seq TLRTWLCCAGSWA/VE

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 555:

Met Thr Ala Gly Thr Leu Arg Thr Trp Leu Cys Cys Ala Gly Ser Trp
-15 -10 -5

Ala Val Glu Leu Pro Ala Glu Pro Leu Val Val Phe Cys Xaa Ser Thr $1 \hspace{1cm} 5 \hspace{1cm} 10 \hspace{1cm} 15$

Ser Arg Lys Arg Ala Lys Gly Leu Ile Gln Ser Val 20 25

- (2) INFORMATION FOR SEQ ID NO: 556:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 26 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi-) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -24..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.3

seq RLLVILCVSVKAG/ST

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 556:

Met Leu Gly Arg Pro Cys Phe His Ser Pro Gln Arg Leu Leu Val Ile -20 -15 -10

Leu Cys Val Ser Val Lys Ala Gly Ser Thr

- (2) INFORMATION FOR SEQ ID NO: 557:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 32 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -28..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.2

seq LQFVLPVATQIQQ/EV

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 557:
- Met Asp Glu Ala Arg Asp Asn Ala Cys Asn Asp Met Gly Lys Met Leu
 -25 -20 -15
- Gln Phe Val Leu Pro Val Ala Thr Gln Ile Gln Gln Glu Val Ile Lys -10 -5 1
- (2) INFORMATION FOR SEQ ID NO: 558:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.2

seq LCALGSAPSSMWA/GE

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 558:
- Met Ser Pro Ile Ser Ile Arg Glu Leu Cys Ala Leu Gly Ser Ala Pro -20 -15 -10
- Ser Ser Met Trp Ala Gly Glu
 -5

- (2) INFORMATION FOR SEQ ID NO: 559:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 69 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -13..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.2

seq MTDLLSASPWALT/IV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 559:

Met Thr Asp Leu Leu Ser Ala Ser Pro Trp Ala Leu Thr Ile Val Ser
-10 -5 1

Ser Glu Leu His Leu Ala Pro Ser Met Thr Thr Val Asp Gln Leu Glu
5 10

Ser Gln Val Asp Asn Val Ile Leu Gln Thr Gly Glu Ser Ala Ser Glu 20 25 30 35

Cys Phe Cys Leu Gln Cys Pro Ser Leu Gly Asn Ile Glu Gly Gly Val 40 45 50

Ala Thr Gly His Xaa 55

- (2) INFORMATION FOR SEQ ID NO: 560:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 70 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -26..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.2

seq LTCGPALVPRLWA/TC

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 560:

Met Ser Trp Ser Gly Leu Leu His Gly Leu Asn Thr Ser Leu Thr Cys
-25 -20 -15

Gly Pro Ala Leu Val Pro Arg Leu Trp Ala Thr Cys Ser Met Ala Thr -10 -5 l 5

Leu Asn Gln Met His Arg Leu Gly Pro Pro Lys Arg Pro Pro Arg Lys
10 15 20

Leu Gly Pro Thr Glu Gly Arg Pro Gln Leu Lys Gly Val Val Leu Cys
25 30 35

Thr Phe Thr Arg Asn Arg
40

- (2) INFORMATION FOR SEQ ID NO: 561:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 67 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -23..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.1 seq LEAFSQAISAIQA/LR
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 561:

Met Ala Asp Val Ile Asn Val Ser Val Asn Leu Glu Ala Phe Ser Gln
-20 -15 -10

Ala Ile Ser Ala Ile Gln Ala Leu Arg Ser Ser Val Ser Arg Val Phe -5 1 5

Asp Cys Leu Lys Asp Gly Met Arg Asn Lys Glu Thr Leu Glu Gly Arg 10 20 25

Glu Lys Ala Phe Ile Ala His Phe Gln Asp Asn Leu His Ser Val Asn $30 \hspace{1cm} 35 \hspace{1cm} 40$

Arg Asp Pro

(2) INFORMATION FOR SEQ ID NO: 562:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 77 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -32..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.1

seg RLLSSLLLTMSNN/NP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 562:

Met Asn Val Ile Asp His Val Arg Asp Met Ala Ala Gly Leu His -30 -25 -20

Ser Asn Val Arg Leu Leu Ser Ser Leu Leu Leu Thr Met Ser Asn Asn -15 -10 -5

Asn Pro Glu Leu Phe Ser Pro Pro Gln Lys Tyr Gln Leu Leu Val Tyr
1 5 10 15

His Ala Asp Ser Leu Phe His Asp Lys Glu Tyr Arg Asn Ala Val Ser 20 25 30

Lys Tyr Thr Met Ala Leu Gln Gln Lys Lys Ala Leu Ser 35 40 45

- (2) INFORMATION FOR SEQ ID NO: 563:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 24 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (3) LOCATION: -16..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4.1

seq ACLAWTAVRPSAC/CH

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 563:

PCT/IB98/01232 WO 99/06550 512

Met Thr Ser Ala Cys Leu Ala Trp Thr Ala Val Arg Pro Ser Ala Cys -15

Cys His Pro Gln Ser Ala Asn Trp 5

- (2) INFORMATION FOR SEQ ID NO: 564:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 77 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -55...1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4 seq VFGMSSSSGASNS/AP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 564:

Met Asn Gly Ser Arg Thr Leu Thr His Ser Ile Ser Asp Gly Gln Leu -45

Gln Gly Gly Gln Ser Asn Ser Glu Leu Phe Gln Gln Glu Xaa Gln Thr -30

Ala Pro Ala Gln Val Pro Gln Gly Phe Asn Val Phe Gly Met Ser Ser

Ser Ser Gly Ala Ser Asn Ser Ala Pro His Leu Gly Phe His Leu Gly

Ser Lys Gly Thr Ser Ser Leu Ser Gln Gln Thr Pro Gly 15

- (2) INFORMATION FOR SEQ ID NO: 565:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 47 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens

(F) TISSUE TYPE: Normal prostate

- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -16..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4

seq FFLFLSFVLMYDG/LR

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 565:

Met Leu Gly Phe Phe Leu Phe Leu Ser Phe Val Leu Met Tyr Asp Gly -15 -10 -5

Leu Arg Leu Phe Gly Ile Leu Ser Thr Cys Arg Val His His Thr Met
1 10 15

Asn Gln Phe Leu Ile Asp Ile Ser Ser Phe Thr Ser Arg Val Arg
20 25 30

- (2) INFORMATION FOR SEQ ID NO: 566:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 48 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -27..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 4

seq SIKVLLQSALSLG/RS

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 566:

Met Met Glu Glu Arg Ala Asn Leu Met His Met Met Lys Leu Ser Ile
-25 -20 -15

Lys Val Leu Leu Gln Ser Ala Leu Ser Leu Gly Arg Ser Leu Asp Ala
-10 -5 1 5

Asp His Ala Pro Leu Gln Gln Phe Phe Val Val Met Glu His Cys Ser 10 15 20

- (2) INFORMATION FOR SEQ ID NO: 567:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 80 amino acids

- (B) TYPE: AMINO ACID(D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -17..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.9

seq XIVSAALLAFVQT/HL

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 567:
- Met Glu Leu Glu Xaa Ile Val Ser Ala Ala Leu Leu Ala Phe Val Gln
 -15 -10 -5
- Thr His Leu Pro Glu Ala Asp Leu Ser Gly Leu Asp Glu Val Ile Phe 1 5 10 15
- Ser Tyr Val Xaa Gly Val Leu Glu Asp Leu Gly Pro Ser Gly Pro Ser 20 25 30
- Glu Glu Asn Phe Asp Met Glu Ala Phe Thr Glu Met Met Glu Ala Xaa 35 40 45
- Val Pro Gly Phe Ala His Ile Pro Arg Gly Thr Ile Gly Xaa Met Met 50 55 60
- (2) INFORMATION FOR SEQ ID NO: 568:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 59 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -26..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.9 seq SLIPLFXFIGTGA/TG
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 568:
- Met Leu Arg Gln Ile Ile Gly Gln Ala Lys Lys His Pro Ser Leu Ile
 -25 -20 -15

PCT/IB98/01232 WO 99/06550 515

Pro Leu Phe Xaa Phe Ile Gly Thr Gly Ala Thr Gly Ala Thr Leu Tyr

Leu Leu Arg Leu Ala Leu Phe Asn Pro Xaa Val Cys Trp Asp Arg Xaa 10 15

Asn Pro Glu Pro Trp Asn Xaa Leu Gly Pro Glu 30

- (2) INFORMATION FOR SEQ ID NO: 569:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 100 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -98..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.8 seq WTSLTCSLVVVDG/CG
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 569:

Met Val Lys Glu Thr Gln Tyr Tyr Asp Ile Leu Gly Val Lys Pro Ser -90

Ala Ser Pro Glu Arg Ser Arg Arg Pro Ile Gly Ser Trp Arg Ser Ser

Thr Thr Arg Thr Arg Thr Arg Met Arg Ala Arg Ser Leu Asn Ser Tyr

Pro Arg His Met Lys Cys Phe Gln Ile Gln Arg Lys Gly Met Phe Met

Thr Lys Ala Glu Ser Arg Gln Xaa Lys Lys Glu Ala Gln Ala Ala Pro

Ala Ser Leu His Pro Tro Thr Ser Leu Thr Cys Ser Leu Val Val Val

Asp Gly Cys Gly

- (2) INFORMATION FOR SEQ ID NO: 570:
 - (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 113 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -36..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.8

seq RALSTXLFGSIRG/AA

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 570:
- Met Ala Asn Leu Phe Ile Arg Lys Met Val Asn Pro Leu Leu Tyr Leu -35 -30 -25
- Ser Arg His Thr Val Lys Pro Arg Ala Leu Ser Thr Xaa Leu Phe Gly -20 -15 -10 -5
- Ser Ile Arg Gly Ala Ala Pro Val Ala Val Glu Pro Gly Ala Ala Val $1 \hspace{1cm} 5 \hspace{1cm} 10$
- Arg Ser Leu Leu Ser Pro Gly Leu Leu Pro His Leu Leu Pro Ala Leu 15 20 25
- Gly Phe Lys Asn Lys Thr Val Leu Lys Lys Arg Cys Lys Asp Cys Tyr 30 35 40
- Leu Val Lys Arg Arg Gly Arg Trp Tyr Val Tyr Cys Lys Thr His Pro
 45 55 60
- Arg His Lys Gln Arg His Met Xaa Thr Leu Ser Leu Gln Ser His Ala 65 70 75

Gln

- (2) INFORMATION FOR SEQ ID NO: 571:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 37 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KET: sig_peptide
 - (B) LOCATION: -32..-1

- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 3.7 seq RIHLCQRSPGSQG/VR
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 571:

Met Ala Ala Ala Ala Ser Arg Gly Xaa Gly Ala Lys Leu Gly Leu
-30 -25 -20

Arg Xaa Ile Arg Ile His Leu Cys Gln Arg Ser Pro Gly Ser Gln Gly
-15 -5

Val Arg Asp Phe Ile

- (2) INFORMATION FOR SEQ ID NO: 572:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 65 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -44..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.7 seq IALTLIPSMLSRA/AG
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 572:

Met Phe Pro Ser Cys Tyr Leu Cys Tyr Ser Leu Cys Gly Ser Ile Leu
-40 -35 -30

Leu Ser Ile Phe Ser Ala Tyr Asn Arg Leu Ser Leu Met Leu Arg Ile
-25 -20 -15

Ala Leu Thr Leu Ile Pro Ser Met Leu Ser Arg Ala Ala Gly Trp Cys
-10 -5 1

Trp Tyr Lys Glu Pro Thr Gln Gln Phe Ser Tyr Leu Cys Leu Pro Cys
5 10 15 20

Gly

- (2) INFORMATION FOR SEQ ID NO: 573:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 65 amino acids

(B) TYPE: AMINO ACID(D) TOPOLOGY: LINEAR

- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -60..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.7

seq QLXFLYFVCCIFQ/DV

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 573:

Met Ser Thr Gln Xaa Gly Leu Ser Met His Ala His Pro Gln Ala Tyr
-60 -55 -50 -45

Thr Pro Phe Ile Tyr Leu His Ala Arg Lys Arg Arg Gly Glu Ile Gly
-40 -35 -30

Asp Ala Asp Ser Arg Phe Asn Asp Arg Tyr Ala His Lys Ser Ala Gln
-25 -20 -15

Leu Xaa Phe Leu Tyr Phe Val Cys Cys Ile Phe Gln Asp Val Tyr Tyr
-10 -5

Xaa 5

- (2) INFORMATION FOR SEQ ID NO: 574:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 39 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -21..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.7 seq SSCSCSLISFTRG/DK
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 574:

Met Lys His Phe Gln Asp Leu Pro Ser Ser Cys Ser Cys Ser Leu Ile
-20 -15 -10

PCT/IB98/01232 WO 99/06550

Ser Phe Thr Arg Gly Asp Lys Tyr Phe Ala Tyr Asn Glu Glu Ile Phe

Leu Val Tyr Asn Ala Asp Gln 15

- (2) INFORMATION FOR SEQ ID NO: 575:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 90 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -62..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.7

seg SILGIISVPLSIG/YC

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 575:
- Met Ser Gln Arg Ser Leu Cys Met Asp Thr Ser Leu Asp Val Tyr Arg ~55
- Xaa Leu Ile Glu Leu Asn Tyr Leu Gly Thr Val Ser Leu Thr Lys Cys
- Val Leu Pro His Met Ile Glu Arg Lys Xaa Xaa Lys Ile Val Thr Val
- Asn Ser I-le Leu Gly Ile Ile Ser Val Pro Leu Ser Ile Gly Tyr Cys
- Ala Ser Xaa His Ala Leu Xaa Gly Phe Phe Asn Xaa Leu Arg Thr Xaa

Leu Ala Thr Tyr Pro Gly Ile Ile Val Ser 25

- (2) INFORMATION FOR SEQ ID NO: 576:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 120 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN

PCT/IB98/01232

- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -98..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.6

seq LALRTSWISSVCS/VT

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 576:

Met Gly Gly Ser Gly Ser Arg Leu Ser Lys Glu Leu Leu Ala Glu Tyr
-95 -90 -85

Gln Asp Leu Thr Phe Leu Thr Lys Gln Glu Ile Leu Leu Ala His Arg
-80 -75 -70

Arg Phe Cys Glu Leu Leu Pro Gln Glu Gln Arg Xaa Xaa Ser Arg His
-65 -55

Phe Gly His Lys Cys Pro Ser Ser Arg Phe Ser Ala Phe Gln Ser Ser -45 -40 -35

Arg Pro Thr Pro Ser Arg Ser Glu Ser Ala Gly Ser Ser Pro His Pro
-30 -25 -20

Gln Pro Lys Thr Ala Leu Ala Leu Arg Thr Ser Trp Ile Ser Ser Val -15 -10 -5

Cys Ser Val Thr Gln Pro Arg Gln Thr Ser Ser Pro Ile Met Pro Ser 1 5 10

Ala Ser Leu Thr Leu Met Met Thr 15 20

- (2) INFORMATION FOR SEQ ID NO: 577:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 51 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -28..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.6 seq PLSDSWALLPASA/GV
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 577:

Met Trp Arg Leu Leu Ala Arg Ala Ser Ala Pro Leu Leu Arg Val Pro
-25 -20 -15

Leu Ser Asp Ser Trp Ala Leu Leu Pro Ala Ser Ala Gly Val Lys Thr -10 -5 l

Leu Leu Pro Val Pro Ser Phe Glu Asp Val Ser Ile Pro Glu Lys Pro 5 10 15 20

Lys Leu Leu

- (2) INFORMATION FOR SEQ ID NO: 578:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 123 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: $-11\overline{4}..-1$
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.6

seq ATFVTQALIQXYA/RI

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 578:
- Met Ala Asp His Val Gln Ser Leu Ala Gln Leu Glu Asn Leu Cys Lys
 -110 -105 -100
- Gln Leu Tyr Glu Thr Thr Asp Thr Xaa Xaa Arg Ser Ser Xaa Ala Glu -95 -90 -85
- Lys Ala Leu Val Glu Phe Thr Asn Ser Pro Asp Cys Leu Ser Lys Cys
 -80 -75 -70
- Gln Leu Leu Glu Arg Gly Ser Ser Ser Tyr Ser Gln Leu Leu Ala -65 -60 -55
- Ala Thr Cys Leu Thr Lys Leu Val Ser Arg Thr Asn Asn Pro Leu Pro
 -50 -45 -40 -35
- Leu Glu Gln Arg Ile Asp Ile Arg Asn Tyr Val Leu Asn Xaa Leu Ala
 -30 -25 -20
- Thr Arg Pro Lys Leu Ala Thr Phe Val Thr Gin Ala Leu Ile Gin Xaa -15 -10 -5
- Tyr Ala Arg Ile Thr Lys Leu Gly Trp Phe Asp

- (2) INFORMATION FOR SEQ ID NO: 579:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 59 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -55..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.6 seq TCSVCCYLFWLIA/IP
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 579:

Met Ala Tyr His Gly Leu Thr Val Pro Leu Ile Val Met Ser Val Phe
-55 -50 -45 -40

Trp Gly Phe Val Gly Phe Leu Val Pro Trp Phe Ile Pro Lys Gly Pro
-35 -30 -25

Asn Arg Gly Val Ile Ile Thr Met Leu Val Thr Cys Ser Val Cys Cys -20 -15 -10

Tyr Leu Phe Trp Leu Ile Ala Ile Pro Ala Trp

- (2) INFORMATION FOR SEQ ID NO: 580:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 128 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -58..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.5 seq GGILMGSFQGTIA/GQ
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 580:

PCT/IB98/01232 WO 99/06550

Met Ser Thr Gly Gln Leu Tyr Arg Met Glu Asp Ile Gly Arg Phe His -55 -50

Ser Gln Gln Pro Gly Ser Leu Thr Pro Ser Ser Pro Thr Val Gly Glu -35

Ile Ile Tyr Asn Asn Thr Arg Asn Thr Leu Gly Trp Ile Gly Gly Ile -20

Leu Met Gly Ser Phe Gln Gly Thr Ile Ala Gly Gln Gly Thr Gly Ala

Thr Ser Ile Ser Glu Leu Cys Lys Gly Gln Glu Leu Glu Pro Ser Gly

Ala Gly Leu Thr Val Ala Pro Pro Gln Ala Val Ser Leu Gln Gly Ser

His Pro Ala Leu Ala Ala Thr Ala Phe Ser Leu Xaa Cys Pro Arg Gly 45

Val Gln Xaa Leu Met Ile Ser Ile Ser Glu His Leu Phe Ile His Ala 65

(2) INFORMATION FOR SEQ ID NO: 581:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 24 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -17..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.5 seq RWWCFHLQAEASA/HP
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 581:

Met Gly Trp Gln Arg Trp Trp Cys Phe His Leu Gln Ala Glu Ala Ser

Ala His Pro Pro Gln Gly Leu Gln 1

(2) INFORMATION FOR SEO ID NO: 582:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 45 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.5 seq VIFFACVVRVRDG/LP
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 582:

Met Ser Val Ile Phe Phe Ala Cys Val Val Arg Val Arg Asp Gly Leu
-15 -5 1

Pro Leu Ser Ala Ser Thr Asp Phe Tyr His Thr Gln Asp Phe Leu Glu
5 10 15

Trp Arg Arg Arg Leu Lys Ser Leu Ala Leu Arg Leu Lys 20 25 30

- (2) INFORMATION FOR SEQ ID NO: 583:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 59 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -16..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 3.5

seq TALAAXTWLGVWG/VR

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 583:
- Met Ala Val Thr Ala Leu Ala Ala Xaa Thr Trp Leu Gly Val Trp Gly -15 -5
- Val Arg Thr Met Gln Ala Arg Gly Phe Gly Ser Asp Gln Ser Glu Asn $\frac{1}{5}$ $\frac{10}{15}$

Val Asp Arg Gly Ala Gly Ser Ile Arg Glu Ala Gly Gly Ala Phe Gly
20 25 30

Xaa Arg Glu Gln Ala Glu Xaa Xaa Arg Tyr Phe 35 40

- (2) INFORMATION FOR SEQ ID NO: 584:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 56 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 12 seq FTLFLALIGGTSG/QY
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 584:

Met Ser Leu Ser Ala Phe Thr Leu Phe Leu Ala Leu Ile Gly Gly Thr -15 -10 -5

Ser Gly Gln Tyr Tyr Asp Tyr Asp Phe Pro Leu Ser Ile Tyr Gly Gln $1 \hspace{1cm} 5 \hspace{1cm} 10$

Ser Ser Pro Asn Cys Ala Pro Glu Cys Asn Cys Pro Glu Ser Tyr Pro 15 20 25 30

Ser Ala Met Tyr Cys Asp Glu Leu - 35

- (2) INFORMATION FOR SEQ ID NO: 585:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1

- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 12 seq FTLFLALIGGTSG/QY
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 585:

Met Ser Leu Ser Ala Phe Thr Leu Phe Leu Ala Leu Ile Gly Gly Thr -15 -10 -5

Ser Gly Gln Tyr Tyr Asp Trp
1 5

- (2) INFORMATION FOR SEQ ID NO: 586:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 96 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -18..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 12

seq FTLFLALIGGTSG/QY

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 586:
- Met Ser Leu Ser Ala Phe Thr Leu Phe Leu Ala Leu Ile Gly Gly Thr
- Ser Gly Gin Tyr Tyr Asp Tyr Asp Phe Pro Leu Ser Ile Tyr Gly Gln $1 \hspace{1cm} 5 \hspace{1cm} 10$
- Ser Ser Pro Asn Cys Ala Pro Glu Cys Asn Cys Pro Glu Ser Tyr Pro
 15 20 25 30
- Ser Ala Met Tyr Cys Asp Glu Leu Lys Leu Lys Ser Val Pro Met Val 35 40 45
- Pro Pro Gly Ile Lys Tyr Leu Tyr Leu Arg Asn Asn Gln Ile Asp His 50 55 60
- Ile Asp Glu Lys Ala Phe Glu Asn Val Thr Asp Leu Gln Trp Leu Gly 65 70 75
- (2) INFORMATION FOR SEQ ID NO: 587:
 - (1) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 111 amino acids
- (B) TYPE: AMINO ACID
- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -20..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 8.9

seq LLLLLLPFLLYMA/AP

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 587:

Met Val Glu Leu Met Phe Pro Leu Leu Leu Leu Leu Leu Pro Phe Leu
-20 -15 -10 -5

Leu Tyr Met Ala Ala Pro Gln Ile Arg Lys Met Leu Ser Ser Gly Val $1 \hspace{1cm} 5 \hspace{1cm} 10$

Cys Thr Ser Thr Val Gln Leu Pro Gly Lys Val Val Val Thr Gly
15 20 25

Ala Asn Thr Gly Ile Gly Lys Glu Thr Ala Lys Glu Leu Ala Gln Arg 30 35 40

Gly Ala Arg Val Tyr Xaa Ala Xaa Xaa Asp Val Glu Lys Gly Glu Leu 45 50 55 60

Val Ala Xaa Glu Ile Gln Thr Thr Gly Xaa Xaa Gln Val Leu Val 65 70 75

Arg Xaa Leu Asp Leu Ser Asp Thr Lys Ser Ile Arg Ala Phe Ala 80 85 90

- (2) INFORMATION FOR SEQ ID NO: 588:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 41 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (C) OTHER INFORMATION: score 8.1

seq LLYLLVPALFCRA/GG

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 588:

Met Trp Leu Leu Tyr Leu Leu Val Pro Ala Leu Phe Cys Arg Ala Gly -15 -5 1

Gly Ser Ile Pro Ile Pro Gln Lys Leu Phe Gly Glu Val Thr Ser Pro 5 10 15

Leu Phe Pro Lys Pro Tyr Pro Asn Gly

- (2) INFORMATION FOR SEQ ID NO: 589:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 71 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -32..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 7.7

seq LLFLVAGLLPSFP/AN

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 589:
- Met Lys Gln Ile Leu His Pro Ala Leu Glu Thr Thr Ala Met Thr Leu -30 -25 -20
- Phe Pro Val Leu Leu Phe Leu Val Ala Gly Leu Leu Pro Ser Phe Pro -15 -5
- Ala Asn Glu Asp Lys Asp Pro Ala Phe Thr Ala Leu Leu Thr Thr Gln 1 5 10 15
- Thr Gln Val Gln Arg Glu Ile Val Asn Lys His Asn Glu Leu Arg Arg 20 25 30

Ala Val Ser Pro Pro Ala Lys 35

- (2) INFORMATION FOR SEQ ID NO: 590:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 138 amino acids
 - (B) TYPE: AMINO ACID

- (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Cancerous prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -17..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 6.9

seq LFLTMLTLALVKS/QD

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 590:

Met Leu Lys Ala Leu Phe Leu Thr Met Leu Thr Leu Ala Leu Val Lys
-15 -10 -5

Ser Gln Asp Thr Glu Glu Thr Ile Thr Tyr Thr Gln Cys Thr Asp Gly
1 5 10 15

Tyr Glu Trp Asp Pro Val Arg Gln Gln Cys Lys Asp Ile Asp Glu Cys
20 25 30

Asp Ile Val Pro Asp Ala Cys Lys Gly Gly Met Lys Cys Val Asn His 35 40 45

Tyr Gly Gly Tyr Leu Cys Leu Pro Lys Thr Ala Gln Ile Ile Val Asn 50 55 60

Asn Glu Gln Pro Gln Gln Glu Thr Gln Pro Ala Glu Gly Thr Ser Gly 65 70 75

Ala Thr Thr Gly Val Val Ala Ala Xaa Ser Met Ala Thr Ser Xaa Val 80 90 95

Leu Xaa Gly Gly Gly Phe Val Ala Ser Ala Ala Ala Val Ala Gly Pro 100 105 110

Glu Met Gln Thr Gly Arg Asn Asn Phe Val

- (2) INFORMATION FOR SEQ ID NO: 591:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 69 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE: -
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
 - (ix) FEATURE:

- (A) NAME/KEY: sig_peptide
- (B) LOCATION: -22..-1
- (C) IDENTIFICATION METHOD: Von Heijne matrix
- (D) OTHER INFORMATION: score 5.9 seq LLILWFHLDCVSS/IL
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 591:

Met Glu Lys Asn Pro Leu Ala Ala Pro Leu Leu Ile Leu Trp Phe His -20 -15 -10

Leu Asp Cys Val Ser Ser Ile Leu Asn Val Glu Gln Ser Pro Gln Ser -5 1 5 10

Leu His Val Gln Glu Gly Asp Ser Thr Asn Phe Thr Cys Ser Phe Pro \$15\$

Ser Ser Asn Phe Tyr Ala Leu His Trp Tyr Arg Trp Glu Thr Ala Lys $30 \hspace{1cm} 35 \hspace{1cm} 40$

Ser Pro Glu Ala Val 45

- (2) INFORMATION FOR SEQ ID NO: 592:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 37 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
 - (ii) MOLECULE TYPE: PROTEIN
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Hypertrophic prostate
 - (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -15..-1
 - (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.3 seq VVTIVILLCFCKA/AE
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 592:

Met Arg Val Val Thr Ile Val Ile Leu Leu Cys Phe Cys Lys Ala Ala -15 -5 1

Glu Leu Arg Lys Ala Ser Pro Gly Ser Val Arg Ser Arg Val Asn His
5 10 15

Gly Arg Ala Gly Gly 20

(2) IMFORMATION FOR SEQ ID NO: 593:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 102 amino acids
 - (B) TYPE: AMINO ACID
 - (D) TOPOLOGY: LINEAR
- (ii) MOLECULE TYPE: PROTEIN
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo Sapiens
 - (F) TISSUE TYPE: Normal prostate
- (ix) FEATURE:
 - (A) NAME/KEY: sig_peptide
 - (B) LOCATION: -90..-1
 - . (C) IDENTIFICATION METHOD: Von Heijne matrix
 - (D) OTHER INFORMATION: score 5.1

seq LLFVATLPFWTHY/LI

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 593:

Met Asp Gln Phe Pro Glu Ser Val Thr Glu Asn Phe Glu Tyr Asp Asp
-90
-85
-80
-75

Leu Ala Glu Ala Cys Tyr Ile Gly Asp Ile Val Val Phe Gly Thr Val -70 -65 -60

Phe Leu Ser Ile Phe Tyr Ser Val Ile Phe Ala Ile Gly Leu Val Gly
-55 -50 -45

Asn Leu Leu Val Val Phe Ala Leu Thr Asn Ser Lys Lys Pro Lys Ser -40 -35 -30

Val Thr Asp Ile Tyr Leu Leu Asn Leu Ala Leu Ser Asp Leu Leu Phe -25 -15

Val Ala Thr Leu Pro Phe Trp Thr His Tyr Leu Ile Asn Glu Lys Gly -10 -5 1 5

Leu His Asn Ala Met Cys - 10

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